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(54) Title: SYSTEMATIC POLYPEPTIDE EVOLUTION BY REVERSE TRANSLATION

(57) Abstract

A method for preparing polypoptide ligands of target molecules wherein candidate mixtures comprised of ribosome complexes or mRNA-polypeptide copolymers are partitioned relative to their affinity to the target and amplified to create a new candidate mixture enriched in ribosome complexes or mRNA-polypeptide copolymers with an affinity to the target.

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SYSTEMATIC POLYPEPTIDE EVOLUTION

BY REVERSE TRANSLATION

FIELD OF THE INVENTION

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target molecule. A method is presented for selecting a acronym for Systematic Polypeptide Evolution by Reverse invention are useful for any purpose to which a binding agents and the like. In addition, polypeptide products hormones, receptors and cell surfaces, nucleic acids, polypeptide ligand that specifically binds any desired diagnostic procedures, cell sorting, as inhibitors of of the invention can have catalytic activity. Target target molecule function, as probes, as sequestering useful to isolate a polypeptide ligand for a desired Translation. The method of the invention (SPERT) is polypeptide ligands that specifically bind a desired including proteins, polysaccharides, glycoproteins, reaction may be put, for example in assay methods, target molecule. The polypeptide products of the molecules include natural and synthetic polymers, target molecule. The method is termed SPERT, an and small molecules such as drugs, metabolites, cofactors, transition state analogs and toxins. We describe herein novel high-affinity

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BACKGROUND OF THE INVENTION

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(1986) Science 232:348). Antigen-encoding mRNAs have Gilmore (1986) J. Cell Biol. 103:2253; Perara et al. These complexes are made of "ribosome complexes" herein, such complexes can be polypeptide encoded by the messenger RNA. Termed isolated by various known processes (Connolly and As translation of mRNA proceeds, stable ribosomes bound to mRNA with tRNA and nascent complexes are formed.

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the partitioning of ribosome complexes according to the present invention is not restricted to immunoreactivity possible. Historically, mRNA selection is closely tied 8.10). Such immunoreactive ribosome complexes can be to immunopurification of ribosome complexes, however, with ribosome complexes (Sambrook, J., Fritsch, E.F., 256, 1495). Cyclical selection and amplification of Maniatis, T. Molecular Cloning: A Laboratory Manual ribosome complexes (Schutz et al. (1977) Nuc. Acids immunoreactivity of nascent polypeptides associated (cold spring Harbor, NY) (1989) <u>ibid</u>. sections 8.9-Res. 4, 71; Shapiro and Young (1981) J. Biol. Chem. protein A column chromatography from non-reactive immunoprecipitated from solution or separated by RNAS with partitionable properties is now also been purified by taking advantage of the of the nascent polypeptides.

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SUMMARY OF THE INVENTION

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Ultimately, both the desired optimal polypeptide ligand partitioned according to their property of binding to a systematic polypeptide evolution by reverse translation carried out in such a way, herein described, that each polypeptide is partitioned together with the means for having a randomized amino acid sequence. Each member (SPERT) includes a candidate mixture of polypeptides given desired target molecule. The partitioning is mRNA encoding a polypeptide is partitioned exactly of the mixture is linked to an individualized mRNA together with that polypeptide. In this way each polypeptide are simultaneously selected, allowing of the desired target and the mRNA encoding the which encodes the amino acid sequence of that further amplifying it by an <u>in vitro</u> process. In its broadest aspect, the method of polypeptide. The candidate polypeptides are

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further synthesis of the selected polypeptide as desired, and further amplification of the coding sequence. It is therefore not necessary to analyze the amino acid sequence of the selected polypeptide (using protein chemistry) in order to produce it in desired quantities.

have taught a method for selective evolution of nucleic mixture. In previously filed applications, U.S. Serial and amplification can be a powerful tool for developing novel compounds when coupled with a partitioning system No. 07/536,428, filed June 11, 1990 and U.S. Serial No. incorporated herein by reference, the inventors herein acids themselves. The insight that cyclical selection is herein adapted to evolving specific coding nucleic selective evolution of a nucleic acid that encodes a polypeptide ligand of a desired target. The present scids based upon binding properties of the nucleic method is therefore a selection based upon coding 07/714,131 filed July 10, 1991, both of which are properties available in a candidate nucleic acid Viewed another way, the invention is the polypeptide ligands binding to target molecules. acids, based on the partitioning properties of

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More specifically, the invention includes a method for making a polypeptide ligand of a desired target molecule which includes the following steps: First, synthesizing a mixture of translatable mRNA's, having certain sequence segments in common such as a ribosome binding site and a translation initiation codon and having a segment encoding a polypeptide at least part of which coding region is a randomized sequence. Second, employing the mRNA mixture in an in vitro translation system. Synthesis of nascent polypeptides ensues, each encoded by its own mRNA. At any time during translation, stable ribosome complexes

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target pairs is thereby enriched for those polypeptides (and, necessarily their coding mRNA's) that can bind to polypeptide ligands. These cycles can be reiterated as binding of each nascent polypeptide to a desired target to be translated mRNA. Although its chemical structure many times as desired, until a desired binding affinity cloned coding mRNA, or by chemical or enzymatic methods molecule. Some polypeptides bind weakly, some tightly, the target. Fourth, the encoding mRNA's are separated ribosome complexes are partitioned with respect to the from the complexes and amplified by conventional means can be isolated. It is preferred to isolate complexes in which translation has been stopped, or "stalled" by is unaltered, translated mRNA is bound to the ribosome nascent peptide and the coding mRNA which is now said complex in a different manner than it was bound prior however conducted, generally separates the mixture of ribosome complexes into ribosome complex-target pairs polypeptide ligand can then be prepared in vivo from and unbound complexes. The set of ribosome complexribosome complex includes at least one ribosome, one to translation, as is known in the art. Third, the some not at all, with the target. The partitioning, polypeptide selected in the foregoing manner can be any of several known circumstances. Each isolated This amplification sets the stage for a subsequent transcription and polymerase chain reaction (PCR). round of transcription, polypeptide synthesis and partitioning to further enrich for target-binding is achieved, or no further improvement in binding cloned and sequenced, if desired. An individual affinity is observed. The coding mRNA for any for amplifying nucleic acids, such as reverse ហ 12 2 20 25 30

invention, means for linking the nascent polypeptide to fixed sequence in the translated region that encodes a freed to translate additional mRNA species. The fewer specific example of this embodiment, a biotin molecule is covalently bound to the 5' end of the mRNA sequence copolymers. By removing the relatively large ribosome affinity interactions--between the polypeptide and the greatly increased. In addition, the ribosome is then the translated mRNA are included in the design of the connection -- either via covalent bonding or very tight mRNA allows for the removal of the ribosomal linkage polypeptides can be generated in the process. In a from the mRNA polypeptide copolymer, the ability to ribosomes that can be utilized, the more randomized between these two elements leaving mRNA.polypeptide partition polypeptides based on the affinity of the polypeptide that may be covalently bound to biotin. utilized, and the nucleic acid template includes a In an alternate embodiment of the present randomized polypeptides to a given target may be system. According to this method, a direct

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a given target molecule. The invention is based on the amplification of nucleic acids can be applied to coding therefore be applied for the first time to up to about Each compound of the invention is a specific ligand of specifically to a desired target compound or molecule. sufficient capacity for forming a variety of two- and products which are polypeptides, each having a unique three-dimensional structures and sufficient chemical sequence, each of which has the property of binding according to the binding affinities of the encoded polypeptides. In vitro evolutionary selection can The present invention provides a class of sequences by partitioning such coding sequences 1018 different polypeptides. Polypeptides have unique insight that cyclical selection and

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physiological limits. For other uses different binding commonly, and preferably, for therapeutic applications, any chemical compound, whether monomeric or polymeric. binding takes place in aqueous solution at conditions versatility available within their monomers to act as ligands (form specific binding pairs) with virtually Molecules of any size can serve as targets. Most of salt, temperature and pH near acceptable conditions can be employed.

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any desired target. The method involves selection from generally applicable to make a polypeptide ligand for The invention also provides a method which is a mixture of candidates and step-wise iterations of selection theme, to achieve virtually any desired structural improvement, using the same general criterion of binding affinity and selectivity.

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possible sequences and structures there is a wide range While not bound by a theory of operation, SPERT polypeptide mixture comprising, for example a 10 amino possibilities. Those which have the higher affinity constants for the target are most likely to bind. is based on the inventors' insight that within a polypeptide mixture containing a large number of acid randomized segment can have 20^{10} candidate of binding affinities for a given target. A

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Additional rounds of SPERT progressively favor the best second polypeptide mixture is generated by translation, predominantly composed of only one or a few sequences. These can then be individually synthesized and tested reverse transcription/amplification/ transcription, a mRNA.polypeptide copolymers, dissociation of mRNA and enriched for the higher binding affinity candidates. for binding affinity as pure ligands. One cycle of ligands until the resulting polypeptide mixture is SPERT effectively achieves reverse translation, at After partitioning ribosome complexes or least quantitatively.

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or family of sequences from a huge number of candidates (referred to herein, along with U.S. Patent Application nucleic acid ligands to a variety of targets -- including times at a moderate stringency it is possible to obtain The ability to rapidly select a single sequence mathematical analysis dramatically demonstrated that by mathematical analysis of the partitioning and cycling has been dramatically shown in the nucleic acid area. acids and protein targets that are not known to bind cycling through the partitioning process a number of the individual species in a randomized mixture which both protein targets that are known to bind nucleic Serial No. 07/536,428, as the SELEX Applications), have the highest affinity to the selected target. In U.S. Patent Application Serial No. 07/714,131 aspects of SELEX referred to as SELEXION. This nucleic acids -- have been identified. In such application there is also a description of a

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In actual practice, the SELEX Applications show similar affinity to the target. The family of ligands was shown to generally have the same three dimensional process by screening a mixture containing a very large that although in some cases a single solution nucleic protein binds to in nature. These results emphasize able to identify a ligand solution that had a higher the practicality of "short cutting" the evolutionary acid ligand may be identified, it is more often the nucleic acid binding protein, the SELEX process was affinity to the protein than the sequence that the case that a family of ligands is identified having Surprisingly, in some cases where the target was a configuration and many conserved sequences. number of candidates.

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repeated until a desired goal is achieved. In the most until no significant improvement in binding strength is general case, selection/amplification is continued Cycles of selection and amplification are

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to provide sufficient polypeptide length to insure that method could, in practice, be used to sample about $10^{18}\,$ desired for combining with other functional domains or the randomized sequence is accessible to the target in number of ways including chemical or enzymic synthesis other containers necessary to perform the method. The allow isolation of two sequence variants in a mixture polypeptides of the test mixture include a randomized selection/amplification method is sensitive enough to polypeptides which could be sampled, only a practical limit dictated by the sizes of reaction vessels and The iterative containing at least 65,000 sequence variants. The sequence portion as well as conserved sequences as the ribosome complex or mRNA polypeptide copolymer. Amino acid sequence variants can be produced in a different polypeptide species. There is no upper limit, in principle, to the number of different of randomized nucleic acid coding sequences. achieved on repetition of the cycle.

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subportions of conserved sequence incorporated with randomized sequence. Sequence variation in coding nucleic acids can be introduced or increased by partially random sequence; it may also contain variable sequence portion may contain fully or mutagenesis before or during the

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applying SPERT to a mixture of candidates comprising a second randomized sequence. The sequence of the first mixture to isolate a second polypeptide ligand, having selected ligand associated with binding or subportions two sequences (one being the first polypeptide ligand) thereof can be introduced into the randomized portion The SPERT procedure is repeated with this second test of the amino acid sequence of a second test mixture. In the case of a polymeric target, such as a first selected polypeptide sequence combined with a protein, the ligand affinity can be increased by selection/amplification iterations.

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A variant of the walking procedure employs a ligand termed "anchor" which is known to bind to the target molecule at a first binding domain (See Figure 8). This anchor molecule can in principle be any molecule that binds to the target molecule and which can be covalently linked directly or indirectly to a small bridge molecule for which a peptide binding sequence is known. When the target molecule is an enzyme, for example, the anchor molecule can be an inhibitor or substrate of that enzyme. The anchor can also be an antibody or antibody fragment specific for the target. The anchor molecule is covalently linked to the bridge molecule, chosen to bind an oligopeptide of known sequence. A test mixture of candidate

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bind to the target molecule at or near the binding site randomized portion and includes also the known sequence above can then be applied to obtain polypeptide ligands isolate polypeptide ligands which bind at a particular molecule binds the polypeptides to the target molecule procedures could employ selections for binding to the target itself. This method is particularly useful to in the vicinity of the anchor binding site. SPERT is surface of the target molecule adjacent to the anchor target are isolated. Walking procedures as described site within the target molecule. The anchor acts to binding site. Polypeptide ligands which bind to the anchor binding site itself or to another part of the ensure the isolation of polypeptide sequences which then applied to select polypeptides which bind a that binds the bridging molecule. The bridging specificity of binding to the target. Walking polypeptides is then prepared which includes a with increased binding strength or increased

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of the anchor.

Screens, selections or assays to assess the effect of binding of a polypeptide ligand on the function of the target molecule can be readily combined with the SPERT methods. Specifically, screens for inhibition or activation of enzyme activity can be combined with the SPERT methods.

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isolating novel polypeptides which are highly specific

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for binding to a particular target molecule.

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In more specific embodiments, the SPERT method provides a rapid means for isolating and identifying polypeptide ligands which bind to nucleic acids and proteins, including enzymes, receptors, antibodies, and glycoproteins.

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In another aspect, the present invention provides a method for detecting the presence or absence of, and/or measuring the amount of a target molecule in a sample, which method employs a polypeptide ligand which can be isolated by the methods described herein. Detection of the target molecule is mediated by its

antibodies in such detection methods and diagnostics is synthesized in vitro or after cloning, since the method be chemically synthesized since its amino acid sequence can be ascertained readily from the nucleotide sequence needs. Compared to antibodies, SPERT-generated ligands have much greater versatility. Conventional antibodies labeled, for example radiolabeled or enzyme linked, to allow qualitative or quantitative detection, analogous One advantage of polypeptide ligands over conventional amplification, e.g., coding nucleic acids, along with the ligand itself. Alternatively, the polypeptide can proteins. The method is more particularly useful for Another advantage is that the entire SPERT process is detecting proteins which are known to be only weakly invention can be employed in diagnostics in a manner of the invention concomitantly selects the means for carried out in vitro and does not require immunizing ligand need not be as large as an antibody molecule. test animals. Furthermore, the binding affinity of similar to conventional antibody-based diagnostics. of its coding mRNA. A SPERT-generated polypeptide to ELISA and RIA methods. The detection method is particularly useful for target molecules which are produce. Thus, polypeptide ligands of the present antibodies of a desired affinity are difficult to polypeptide ligands can be tailored to the user's binding to a polypeptide ligand specific for that target molecule. The polypeptide ligand can be antigenic, or for which conventional monoclonal that polypeptides are capable of being readily

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Polypeptide ligands of small molecule targets are useful as diagnostic assay reagents and have therapeutic uses as sequestering agents, drug delivery vehicles and modifiers of hormone action. Catalytic polypeptides are selectable products of this invention. For example, by selecting for binding to transition state analogs of an enzyme catalyzed reaction, catalytic polypeptides can be selected. Catalytic immunoglobulins have been developed by raising antibodies to transition state analogs (Schultz, P.C. (1989) Angew. Chem. Int. 2d Engl. 28:1283-1295; Schultz, P.G. (1989) Acc. Chem. Res. 22:287-294; Pollack, S.J. et al. (1989) Meth. Enzymol. 178:551-

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isolated by SPERT. Polypeptide ligands which bind to a useful application of this method is to inhibit protein function, for example to inhibit receptor binding or to to target molecules which are proteins. A particularly target molecule using polypeptide ligands which can be modification. This method is particularly applicable inhibit enzyme catalysis. In this case, an amount of modifying the function of the target is combined with the target molecule to achieve the desired functional for example to select inhibitors or activators of the the selected polypeptide molecule which is effective In yet another aspect, the present invention specifically modify function of the target molecule, target molecule are screened to select those which function of the target molecule. An amount of the selected polypeptide ligand which is effective for or target protein inhibition is combined with the provides a method for modifying the function of a target protein to achieve the desired inhibition.

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are immunoglobulins, which, although capable of a large

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repertoire of binding affinities, are nevertheless

variations of a narrow amino acid sequence and

structural theme. SPERT-generated polypeptide ligands,

in contrast, are unlimited as to structural type, and

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therefore have virtually unlimited potential for

The term "reverse translation" is used throughout as shorthand for the concept of information flow from polypeptide sequence to nucleic acid sequence. The phrase and shorthand make reference to

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the original and revised "central dogma" pronounced by Prancis Crick many years ago. Crick understood and articulated the idea that either RNA or DNA could serve as a template for the synthesis of complementary nucleic acid sequences, and that chemically either RNA or DNA could serve as a template for the synthesis of both RNA and DNA. Crick noted that proteins, comprised of strings of amino acids, were templated by nucleic acid but could not serve themselves as a template for the synthesis of nucleic acids.

Most importantly, no simple chemistry is known that allows "reverse translation"; that was the basis nearly 25 years ago of Crick's adaptor hypothesis for using information in RNA to yield specified protein sequences during translation.

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containing a sequence specified by a polypeptide (whose although binding of a small number of polypeptides from mRNAs that encode those very polypeptides that provides polypeptides to a selected target are known in the art, postulates. While no process, no simple chemistry, is ribosome complex or mRNA polypeptide copolymer of the mechanism for amplifying and using mRNAs that encode sequence is unknown to the scientist at the time of a randomized pool of polypeptides is of no value by SPERT has at its center a form of reverse reverse translation), SPERT provides a reliable known that provides synthesis of a nucleic acid itself. It is the concomitant selection in the translation that does not conflict with Crick's polypeptides of desired function but of unknown sequence. Techniques for binding one or a few a form of reverse translation because:

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 the selected coding sequences can be amplified to yield large quantities of both DNA and RNA:

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 the newly made mRNA can be used for synthesizing polypeptides, now a smaller set than the

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original randomized mixture of polypeptides from which non-binding, or poorly-binding polypeptides have been removed, and;

3) the polypeptides held in ribosome complexes or mPNA.polypeptide copolymers can be used for a subsequent round of SPERT.

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Finally, "reverse translation" during SPERT does not yield a nucleic acid from only polypeptide sequence, but "reverse translation" does provide (through amplification techniques) net synthesis of the templates from which the desired polypeptide was synthesized. In principle a single molecule of polypeptide of the desired activity, along with a single template RNA in the translation complex or copolymer, will lead to a nanomole or even a micromole of nucleic acid corresponding to that polypeptide sequence. This net synthesis of nucleic acids based on the partitioning and activity of the desired polypeptide is an effective quantitative reverse translation that provides the materials for subsequent

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rounds of SPERT.
Also, the coding sequence can be used to deduce the amino acid sequence of a selected polypeptide. The polypeptide can then be synthesized by chemical

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25 methods, if desired.

BRIEF DESCRIPTION OF THE FIGURES

Figure 1 is a diagrammatic representation of steps in the process of the invention. The top panel depicts a double-stranded DNA template having a T7 promoter ("T7 PRO") and a segment of randomized sequence, represented as "nnn.....", preceded by a start codon, ATG. The initiation site of transcription and direction of transcription are shown as a vertical line labeled "+1" and an arrow, respectively. In vitro transcription creates mRNAs (2nd panel) which contain, from left to right, a ribosome binding site, a

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polypeptide is depicted as a thick white line extending The region bordered by a dotted line is expanded in the Figure 2 is a diagram showing expanded views of complex as in the third panel of Figure 1. A cut-away acids of the nascent polypeptide buried in the complex and unavailable for interaction with the solvent. The The carboxy-terminal end of the nascent polypeptide is shown connected to a peptidyl-tRNA (curly black line). covalently linked to a transfer RNA molecule which is hydrogen-bonded to the mRNA at a codon in the P-site. bottom panel showing that the nascent polypeptide is ribosome is depicted as 30-40 amino acids in length. view of the ribosome (2nd panel) shows 30-40 amino vertically from a central tunnel (black) near the center of the ribosome. That portion inside the a ribosome complex. The top panel is a ribosome ribosome is depicted with two shades of gray to indicate inner and outer regions. The nascent

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Figure 3 is a diagram that represents partitioning polypeptide ligands by direct immunoprecipitation. The top panel is a ribosome complex as in Figure 1. The center panel depicts several ribosome complexes where the nascent polypeptide is represented as a short, thick white line with hatching to indicate the segment of randomized sequence. Molecules of a first antibody

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(immunoglobulin) are represented as inverted Y-shaped structures drawn with heavy, straight black lines.
Interaction (binding) of a nascent polypeptide with the epitope recognition site of an immunoglobulin is shown for two ribosome complexes. Nascent polypeptides are selected that have affinity for immunoglobulin molecules. The bottom panel shows addition of a second antibody (white inverted Y's) generally reactive to the first immunoglobulin resulting in an immunoprecipitate containing the selected ribosome complexes, shown as a cluster in the left half of the panel.

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for the target protein are shown in white. Those with affinity for the "pan" are shown in light gray labeled associated with the target protein, through its "pan", depicted in the second panel. Those with no affinity with affinity for the "handle" are dark gray, labeled directed against the "handle" either displaces ligand "handle". These "P" complexes are immunoprecipitated Figure 4 is a diagram showing partitioning of immunoreactive domain ("handle") and a target domain attached by the "handle" to the nascent peptide. In with the "P" complexes made up of a ribosome complex polypeptide ligands by indirect immunoprecipitation. attached by the "pan" to the nascent peptide. Those associations of the "H" complexes or those complexes are unreactive. The first antisera form a sandwich The top panel shows a target protein which has an with an "H" and shown with a bound target protein and bound to the first immunoglobulin through the with a "P" and shown with a bound target protein ("pan"). Three types of ribosome complexes are by second antisera directed against the primary the third panel, a first antibody (black lines) antisera, as shown in the bottom panel.

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Figure 5 is a diagram showing selection of polypeptide ligands by membrane partitioning. The top panel shows a ribosome complex as in Figure 1. The

middle panel shows ribosome complexes and membrane vesicles with membrane proteins. The membrane vesicles are depicted as a hatched band interrupted by hatched ovals that depict membrane proteins embedded in the membrane. In the middle panel, ribosome complexes are shown binding with membrane protein so that the nascent polypeptides having binding affinity for a membrane protein are partitioned. The bottom panel depicts three ribosome complexes bound to a membrane vesicle, forming a large complex which is separable from unbound ribosome complexes.

Figure 6 is a diagram showing partitioning of polypeptide ligands by affinity column chromatography. Ribosome complexes (top panel) are passed through a column containing insoluble support materials to which have been bonded target molecules. The middle panel is an expanded view of the column showing support materials (hatched circular segments) with attached target molecules (black bars) to which some ribosome complexes are bound. The bottom panel shows, enlarged, a single ribosome complex in which the nascent polypeptide (light shading) is bound to a target molecule which is attached to a column support bead (hatched). Ribosome complexes with high affinity to the target molecules are retained on the column and subsequently eluted to continue with SPERT.

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Figure 7 is a diagram showing anchoring of a binding epitope and secondary ligand evolution. A molecule ("inhibitor") of known affinity for a target site on a protein is covalently linked to a "guide epitope". The guide epitope is any molecule for which there exists a peptide ligand, including a portion of a monoclonal antibody which contains an epitope recognition domain (Fab fragment). The mRNA encodes a reactive peptide sequence that binds the guide epitope, incorporated into the nascent polypeptide. The bottom panel depicts a ribosome complex having a nascent

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polypeptide that includes the reactive, guide binding, segment (shaded) and a randomized segment (unshaded). The ribosome complex is shown bound to the protein of interest by a binding interaction between the guide epitope and the reactive segment and by a secondary binding interaction between the randomized segment and a neighboring site on the target protein of interest. The randomized portion of the nascent polypeptide is free to evolve interactions with secondary sites on the target protein.

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Figure 8 is a diagram which shows the DNA to be transcribed and the relationships of the oligonucleotides of Tables 1 and 2 in the DNA, prior to inserting the randomized sequence. The depicted structure constitutes a cassette for carrying out the transcription, translation, reverse transcription and pCR processes used in SPERF.

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DETAILED DESCRIPTION OF THE INVENTION

The following terms are used herein according to the definitions.

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polypeptide is used herein to denote any string of amino acid monomers capable of being synthesized by an <u>in vitro</u> translation system. The term also embraces post-translational modifications introduced by chemical or enzyme-catalyzed reactions, as are known in the art. Such post-translational modifications can be introduced prior to partitioning, if desired. Unless specified herein, all amino acids will be in the L-stereoisomeric form. Amino acid analogs can be employed instead of the 20 naturally-occurring amino acids. Any amino acid analog that is recognized by an aminoacyl-tRNA synthetase can be employed. Several such analogs are known, including fluorophenylalanine, norleucine, azetidine-2-carboxylic acid, S-aminoethyl cysteine, 4-

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azetidine-2-carboxylic acid, S-aminoethyl cysteine, 4 methyl tryptophan and the like. Ligand means a polypeptide that binds another

ligand. The candidate mixture of nucleic acids serving as source of a candidate mixture of polypeptides can be nucleic acids or nucleic acids made by a combination of Candidate mixture is a mixture of nucleic acids in vitro transcription products of naturally-occurring and of polypeptides of differing sequence, from which compound of interest for which a ligand is desired. A cofactor, inhibitor, drug, controlled substance, dye, to select a desired coding seguence and/or a desired component, antigen, antibody, virus, virus component, synthesized nucleic acids, enzymatically synthesized the foregoing techniques. Target molecule means any protein, carbohydrate, polysaccharide, glycoprotein, peptide, enzyme, nucleic acid, nucleic acid binding nutrient, growth factor, toxin, lipid, glycolipid, hormone, receptor, receptor ligand, cell membrane target molecule can be a protein, fusion protein, substrate, metabolite, transition state analog, nucleic acids or fragments thereof, chemically etc., without limitation.

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Partitioning means any process whereby ribosome complexes or mRNA.polypeptide copolymers bound to target molecules, termed complex-target pairs herein, can be separated from ribosome complexes or mRNA.polypeptide copolymers not bound to target molecules. Partitioning can be accomplished by various methods known in the art. The only requirement is a means to separate complex-target pairs from unbound ribosome complexes or mRNA.polypeptide copolymers.

Columns which selectively bind complex-target pairs but

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not ribosome complexes or mRNA.polypeptide copolymers, (or specifically retain ligand to an immobilized target) can be used for partitioning. A membrane or membrane fragment having the target on its surface can bind ligand-bearing ribosome complexes or mRNA.polypeptide copolymers forming the basis of a partitioning based on particle size. The choice of partitioning method will depend on properties of the target and of the complex-target pairs and can be made according to principles and properties known to those of ordinary skill in the art.

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copies of a molecule or class of molecules. Amplifying proportions of different sequences in the mixture prior reaction to increase the copy number of each cDNA, and transcribing the cDNA copies to obtain an abundance of Amplifying means any process or combination of reactions known in the art can be used as appropriate, cDNA copies of selected mRNAs, using polymerase chain carried out by a sequence of three reactions: making process steps that increases the amount or number of amplification and the like, as will be recognized by those skilled in the art. The amplification method coding mRNA molecules in the disclosed examples is should result in the proportions of the amplified mRNA molecules having the same sequences as the selected mRNAs. Any reaction or combination of mixture being essentially representative of the including direct DNA replication, direct mRNA to amplification.

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Specific binding is a term which is defined on a case-by-case basis. In the context of a given interaction between a given ligand and a given target, a binding interaction of ligand and target of higher affinity than that measured between the target and the candidate ligand mixture is observed. In order to compare binding affinities, the conditions of both binding reactions must be the same, and should be

that demand the reguisite specificity during SPERT, or comparable to the conditions of the intended use. For made that reflect the interaction between ligand as a whole and target as a whole. The polypeptide ligands of the invention can be selected to be as specific as required, either by establishing selection conditions "walking" and other modifications using iterations of the most accurate comparisons, measurements will be by tailoring and modifying the ligands through

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an <u>in vitro</u> translation system can introduce additional Randomized is a term used to describe a segment any possible sequence over a given length. Randomized of the genetic code, and biases in the tRNA content of Introducing a deliberate bias into a randomized coding region can reduce the bias of the resulting translated instead of "random" to reflect the possibility of such of a nucleic acid or polypeptide having, in principle nucleic acid sequences will be of various lengths, as which random sequence segments are made may not yield mathematically random sequences due to unknown biases or nucleotide preferences that may exist. Redundancy nucleotides. The chemical or enzymatic reactions by desired, ranging from about twelve to more than 300 amino acid sequence. The term "randomized" is used synthesis, large deviations are not known to occur. deviations from non-ideality. In the techniques presently known, for example sequential chemical bias in the translated amino acid sequences.

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randomized sequence, for example, by altering the molar polypeptides or to lower the frequency of appearance of triphosphates of the synthesis reaction. A deliberate A bias may be deliberately introduced into a ratios of precursor nucleoside (or deoxynucleoside) bias may be desired, for example, to improve the randomness of amino acid sequence of translated certain amino acids.

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biased for the type of structure likely to bind a given or polar (Ser). Randomized sequences biased for codons target. For example, polypeptide sequences biased for of the form GRN are biased for acidic amino acids, Asp in the triplet codon will lack termination signals and will not encode amino acids Phe, Tyr, Cys and Trp. By commonly encoded amino acids are basic (Arg, Asn, Lys) acidic amino acids can bind cationic target molecules Randomized sequences in which U is never the 1st base For example, a randomized sequence biased for Adenine or Guanine and N is any nucleotide) the most such strategies, randomized coding sequences can be (GAU, GAC) and Glu (GAA, GAG), and Glycine (GGN). codons of the form ARN (where A is Adenine, R is more easily than completely random polypeptides.

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orientation and sequence proximal to the 5' end of the such as UUG and GUG can serve as initiation codons and encode methionine if properly spaced within a ribosome In prokaryotes, as is known in the art, other codons, requisite sequences for translation in a conventional RNA, a ribosome binding site and an initiation codon. Translatable mRNA is RNA which possesses all in vitro translation system. These include, proper binding site.

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the initiation codon. The translation initiation codon art. In procaryotic systems, the ribosome binding site GAGG or AGGA, usually located about 5 - 12 bases 5' to sequence in the mRNA which functions as a binding site is therefore usually located within 5 - 12 bases from is a short purine-rich region with a sequence such as procaryotic or eucaryotic origin, as is known in the the ribosome binding site in the 3' direction on the mRNA. These sequences are sometimes termed a Shinesequences which function as ribosome binding sites for a ribosome in an <u>in vitro</u> translation system. differ depending on whether the ribosomes are of Ribosome binding site means a nucleotide

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Dalgarno sequence. The structures of ribosome binding initiation of protein synthesis are well known in the sites and their proper placement to ensure correct

trinucleotide sequence AUG which encodes methionine and polypeptide and also sets the codon reading frame for the nucleotide sequence in the 3' direction from the which encodes a first amino acid of an encoded Initiation codon is a characteristic initiation codon.

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including at least one ribosome, attached mRNA molecule with attached ribosome(s) and nascent peptide(s) remain and, for each ribosome, a nascent polypeptide attached via tRNA to the ribosome. The nascent polypeptide has Ribosome complexes are formed, as is known in the art, an amino acid sequence encoded by the attached mRNA. Ribosome complex is a macromolecular complex tRNA, etc., as known in the art. The mRNA together termination codon in the message, lack of a charged during protein synthesis. Ribosome complexes are stable if they become stalled for any reason, for example, by depletion of release factor, lack of stably bound and can be isolated together, using methods known in the art.

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non-translated and translated regions so that a portion as well to facilitate the formation of mRNA.polypeptide of the translated polypeptide will link with a portion of a candidate mixture in which the RNA includes fixed mRNA.polypeptide copolymers are formed by the creation mRNA.polypeptide copolymer is a macromolecular complex including an mRNA and a polypeptide having an polypeptides or tRNA species utilized may be modified interaction. In other embodiments, the translated of the mRNA via a covalent bond or tight affinity amino acid sequence encoded by the attached mRNA. sequences and/or chemical modifications in both According to one embodiment of the invention,

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or mRNA.tRNA.polypeptide copolymers.

commercially. The conditions for carrying out in vitro are the $\overline{ extbf{E}_{m{\cdot}}}$ coli system, the wheat germ system, and the In vitro translation can be carried out using rabbit reticulocyte system. The latter is available known systems. These well-known translation systems translations are well-known in the art, and various modifications, adaptations and optimizations are available to those skilled in the art.

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The combination of translatable mRNA encoding a polypeptide and in vitro translation system constitute polypeptide encoded by the mRNA. The mRNA can itself be amplified using reverse transcription, PCR with amplified mRNA can serve for in vitro synthesis of amplifying means for amplifying the quantity of desired quantities of the encoded polypeptide. noted, <u>supra</u>, this process constitutes reverse appropriate primers and an RNA polymerase. translation.

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The terms "ribosome" and "nascent peptide" have "translated mRNA" simply refers to mRNA present in a conventional meanings known in the art. The term ribosome complex, either wholly or partially translated.

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complexes of which the nascent polypeptide component is bound to a target molecule. The target molecule can be free in solution or bound to a solid support matrix. Ribosome complex-target pairs are ribosome

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peptides ABCDX or ABXDE are 80% homologous but peptides Homology is used to compare the related uses of ABXYZ, AXYZE and XYZDE are 40% homologous and peptides occupied by the same amino acid in two polypeptides is position by position. The percent of those positions sequences. Percent amino acid sequence homology is the percent sequence homology. Thus, given peptide ABCDE as a naturally-occurring comparison peptide, measured by comparing sequences of equal length

EDCBA, BDAEC, MNOPQ are non-homologous.

selection of polypeptide ligands which bind to a target Iterative cycling of the selection/ amplification steps molecule, for example a protein, with amplification of The SPERT method involves the combination of a polypeptides which bind most strongly to the target those selected polypeptides via the attached mRNAs. from a pool which contains a very large number of nucleic acids and hence encoded polypeptides. allows selection of one or a small number of

the ultimate case until a single species remains in the in binding over background levels during cycling of the selection/ amplification. In such cases, the sequence achieved, For example, cycling can be continued until test mixtures of polypeptides show limited improvement test mixture is achieved or until a minimum number of polypeptide components of the mixture is obtained (in binding is achieved. It may be the case that certain a desired level of binding of the polypeptides in the increased until improvements in binding are achieved. test mixture). In many cases, it will be desired to Anchoring protocols and/or walking techniques can be and length variation in the test mixture should be continue cycling until no further improvement of procedure is continued until a selected goal is Cycling of the selection/amplification employed as well.

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polypeptides. A translatable mRNA mixture is prepared, each member of the mixture including in its nucleotide sequence a ribosome binding site, an initiation codon individual mRNA's contain a randomized region flanked Specifically, the method requires the initial facilitate amplification of selected nucleic acids. by sequences conserved in all nucleic acids in the and a randomized coding region. Preferably the mixture. The conserved regions are provided to preparation of a test mixture of candidate

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method of amplification. The randomized coding region skill in the art can make, having in mind the desired the choice of sequence is one which those of ordinary Since thère are many such sequences known in the art,

process. In principle, the mRNA's employed in the test variability in the polypeptide test mixture can also be amplified. The method of the present invention is most subportions that are randomized, along with subportions introduced or augmented by generating mutations in the which are held constant in all nucleic acid species in practically employed for selection from a large number selected for binding, to the target can be integrated binding or improved specificity of binding. Sequence code for amino acid sequences that bind, or have been the mixture. For example, sequence regions known to of sequence variants. Thus, it is contemplated that Depending on the desired polypeptide structure, the assess binding of polypeptide sequences ranging in with randomized coding regions to achieve improved the present method will preferably be employed to can have a fully or partially randomized sequence coding mRNA's during the selection/amplification mixture can be any length as long as they can be coding portion of the nucleic acid can contain according to the desired translation product. Ŋ 2 12 20 25

provides the advantages of being precisely controllable thereof) or by synthesis of a template from which the nucleic acid (or portions thereof) can be prepared by as to length and allowing individual randomization at acids in the test mixture can be derived in a number The randomized portion of the coding nucleic chemical synthesis of the nucleic acid (or portions use of appropriate enzymes. Chemical synthesis of ways. For example, full or partial sequence randomization can be readily achieved by direct

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length from about four amino acids to any attainable

size.

generally be preferred that the test mixture contain as Therefore, candidate mRNA mixtures that have randomized sequences for all to be sampled in one selection. Many selected fragments of partially digested (or otherwise mixture. Alternatively, the synthesizer can be set up End addition, catalyzed by terminal transferase in the nucleotide triphosphates can add a randomized seguence sequence is employed, it is not necessary (or possible cleaved) preparations of large, natural nucleic acids, four activated nucleotide substrates or with a biased can be used, either with an equivalent mixture of the to provide a limited nucleotide selection at a given position, e.g., only A at the first triplet position. nucleic acids can also be achieved by employing sizepractical matter, it is possible to sample only about epitotes recognized by antibodies are only 5-10 amino each triplet position. A commercial DNA synthesizer from long randomized segments) that the test mixture 10^{36} different candidate nucleic acid sequences which translated polypeptide are identified. A randomized sequence of 60 nucleotides will contain a calculated presence of nonlimiting concentrations of all four large a number of possible sequence variants as is to a segment. Sequence variability in the coding segments longer than 60 contain too many possible preparations. In those cases in which randomized contains all possible variant sequences. It will practical for selection, to insure that a maximum 10¹⁸ polypeptide candidates in a single selection. such as genomic DNA preparations or cellular RNA number of potential amino acid sequences of the would encode 10^{26} possible decapeptides. As a

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preferred that any conserved regions employed in the test nucleic acids do not contain sequences which interfere with amplification.

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laboratory environment. A system that operates to form higher than the limitations found in other systems that randomized polypeptides. However, when the ribosome is be practically tested at a time rises to at least about $10^{17} \ \text{to} \ 10^{18} \ \text{different candidate sequences, depending on}$ ribosome complexes requires a stoichiometric amount of ribosomes. The limitation of 10^{12} to 10^{14} complexes is ribosome in the translation mixture. The presence of this quantity of ribosomes severely limits the amount ribosomes per cell, a huge amount of E. coli would be reaction mixture, the number of mRNA species that can volume or mass of materials that can be handled in a number of candidates that may be sampled include the of sequences that can be sampled -- to about 1012 to would be impractical. As $\overline{ ext{E. coli}}$ has only about 10^4 the number of mRNAs translated by a single ribosome. not bound up in the ribosome complex but is free to The practical considerations that limit the quanitites of ribosomes in excess of these amounts 10" complexes. The production and isolation of translate a large number of mRNA species in the have been devised for sampling large numbers of required to produce stoichiometric amounts of

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The complex of a ribosome, mRNA, and nascent polypeptide attached to a tRNA in the P-site of the ribosome is very stable. Release of the nascent peptide from the complex and of the mRNA from the ribosome requires protein release factors. Release factor recognition requires the positioning of the stop codons of the mRNA in the A-site of the ribosome. In the absence of a stop codon or release factor the dissociation of the translation complex from mRNA is very slow. The addition of the antibiotics cycloheximide (eukaryotic systems) and chloramphenicol

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possible sequences of a candidate mixture to select a

acids in length. It is not necessary to sample all

polypeptide ligand of the invention. It is basic to the method that the coding nucleic acids of the test mixture are capable of being amplified. Thus, it is

(prokaryotic system) further stabilizes the complexes chromatography and gradient centrifugation can be so that extensive manipulations like column performed.

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analogues, and the use of certain antibiotics will also translation systems which have been depleted of release defined sites is to include at the 3' end of the coding factor (by immunoinactivation or mutation) will result nascent polypeptide for the successful partitioning of mutants under the restrictive condition will result in ways in which this can be accomplished. Because stop translating ribosome that does not encounter any stop paused at the end of a coding sequence on a mRNA with codons will proceed to the end of a mRNA and stall at stall translational complexes. The timed addition of translation reaction can produce predictable sizes of region a stretch of sense codons which are recognized codons. Removal of GTP, the use of non-hydrolyzable by a single species of tRNA for which there exists a stalled complexes at the stretch of sense codons for exist temperature-sensitive tRNA synthetase mutants. partitioning of the complex. There are a number of the 3' end (Connolly and Gilmore, <u>supra</u>). In <u>vitro</u> the translational complex. In some organisms there In this embodiment a ribosome is preferably Another way of stalling translational complexes at codons are essential for release factor action, a these exogenous factors to a synchronous in vitro in the stalling of translation complexes at stop translation reactions done from extracts of such the encoded nascent polypeptide available for conditional tRNA synthetase mutant. In vitro that particular tRNA.

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partitionable ribosome complexes. Stable complexes can be isolated at any time during active translation. It It will be understood that it is not necessary to stall or pause the translation process to obtain

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Ribosome complexes isolated during active translation constitute a population of nascent peptides of varied length. By synchronously initiating translation and length of the randomized segment, e.g., to test the effects of polypeptide length on binding efficacy. thereafter, the effects of increasing polypeptide ribosome complexes when it is desired to vary the is advantageous to isolate actively translating isolating ribosome complexes at various times length can be compared.

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Sci. 85:9436-9440; and in U.S. Patent 4,683,195 (Mullis Polymerase chain reaction (PCR) is an exemplary et al.) and U.S. Patent 4,683,202 (Mullis et al.). In cycles of replication of a desired single-stranded DNA method for amplifying nucleic acids. Descriptions of 233:1076-1078; Innis et al. (1988) Proc. Natl. Acad. its basic form, PCR amplification involves repeated PCR methods are found, for example in Saiki et al. (1985) Science 230:1350-1354; Saiki et al. (1986) Nature 324:163-166; Scharf et al. (1986) Science

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oligonucleotide primers complementary to the 3' ends of and DNA denaturation. Products generated by extension from one primer serve as templates for extension from both strands, primer extension with a DNA polymerase, (or cDNA copy of an RNA) employing specific

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of (Burg et al.) requires the presence or introduction a promoter sequence upstream of the sequence to be described in PCT published application WO 89/01050 amplified, to give a double-stranded intermediate. the other primer. A related amplification method

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them be subject to another round of amplification with polymerase. The resultant RNA copies are treated with RNA polymerase. Alternative methods of amplification stranded promoter containing intermediates which can reverse transcriptase to produce additional double-Multiple RNA copies of the double-stranded promotercontaining intermediate are then produced using RNA

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include among others cloning of selected DNAs or cDNA copies of selected RNAs into an appropriate vector and introduction of that vector into a host organism where the vector and the cloned DNAs are replicated and thus amplified (Guatelli, J.C. et al. (1990) Proc. Natl. Acad. Sci. <u>81</u>:1874). In general, any means that will allow faithful, efficient amplification of selected nucleic acid sequences can be employed in the method of the present invention. It is only necessary that the proportionate representations of sequences after amplification reflect the relative proportions of sequences in the mixture before amplification.

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promoter in their 5' portions. Full-length cDNA copies then transcribed in vitro. Transcripts are used in the transcriptase primed with an oligomer complementary to to the conserved 5' and of the selected RNAs. Doublethe 3' seguences of the selected RNAs. The resultant promoter sequence as well as a sequence complementary Specific embodiments of the present invention next selection/ amplification cycle. The method can designed to contain a sequence transcribed from a T7 for amplifying RNAs are based on Innis et al. (1988) stranded products of this amplification process are supra. The RNA molecules in the test mixture are of selected mRNA molecules are made using reverse cDNAs are amplified by Tag DNA polymerase chain extension, employing a primer containing the T7 optionally include appropriate nucleic acid purification steps.

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In general, any protocol which will allow selection of polypeptides based on their ability to bind specifically to another molecule, i.e., a protein or any target molecule, can be employed in the method of the present invention. It is only necessary that the ribosome complexes or mRNA.polypeptide copolymers be partitioned without disruption such that the selected coding mRNA's are capable of being amplified.

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non-target binding complexes are eluted from the column polypeptide ligands of the target are retained and the and coupling reactions is available for application of will depend on the type of target, the strength of the polypeptides together with mRNA's encoding each remain elution procedures is within the skill of the ordinary chromatography techniques, including support matrices polypeptide employed to achieve the desired selection For example, in a column binding selection in which a protein to test polypeptides in the incubated mixture that any polypeptide that binds to the target will be binding that is present. The relative concentrations needed to achieve the desired partitioning result can available binding sites occurs and those polypeptides with appropriate buffer. A wide variety of affinity bound to the column. The relative concentrations of for. When polypeptide is in excess, competition for when an excess of target is employed, it is expected binding interaction and the level of any background influences the strength of binding that is selected selected. The relative concentrations of target to immobilized target molecules, the complexes bearing experimentation. Similarly, it may be necessary to background binding. Again such optimization of the which bind most strongly are selected. Conversely, test mixture of ribosome complexes bearing nascent randomized polypeptide is passed over a column of optimize the column elution procedure to minimize be readily determined empirically without undue a column partitioning system. Target binding artisan. ល ព 20 15 25 30

An unexpected feature of the invention is the fact that the polypeptide ligand need not be elutable from the target to be selectable. This is because it is the mRNA that is recovered for further amplification or cloning, not the polypeptide itself. It is known that some affinity columns can bind the most avid

into their component units based on the specific nature However the method of the invention can be successfully practiced to yield avid ligands, even covalent binding including ligands that bind tightly, irreversibly or Various mRNA.polypeptide copolymers may be separated amplified, as described elsewhere herein, to yield a ligands so tightly as to be very difficult to elute. sulfate without affecting the integrity of the mRNA. mixture of coding sequences enriched for those that encode polypeptide ligands of the desired target, ligands. Ribosome complexes can be disrupted by polypeptide. The mRNA's of selected ligands are denaturing agents such as urea or sodium dodecyl of linking between the RNA and the associated covalently.

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or cycloheximide which stall translational complexes on Staphylococcus aureus and has a high affinity for IgGs. immunoreactive ribosomes on a protein A column. (IgGs embodiment, ribosome complexes are purified from cells background of mRNAs which do not encode the epitope of complex can be lowered by using purified IgGs against ribosome complexes or mRNA.polypeptide copolymers can in the presence of inhibitors such as chloramphenicol Immunoreactivity of nascent polypeptides on are one class of the soluble immunoglobulins which epitope of interest followed by binding antibodies interest but are trapped by the immunoprecipitated Protein A binding does not interfere with epitope mRNA. Binding of antibodies which recognize the containing the mRNAs which encode the epitope. be used to purify the encoding mRNAs. In one immunoprecipitation of the ribosome complexes compose antisera. Protein A is derived from which recognize those antibodies results in the epitope followed by purification of the recognition.)

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These procedures for immunoprecipitation to

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copolymers can be used in a variety of modifications to Ribosome complexes or mRNA polypeptide copolymers which 4). A protein is composed of an immunoreactive domain for which known antibody exists, and a separate target domain for which one wishes to evolve protein ligands. interact with the target domain (the "pan") via their such modification is termed "panhandling" (See Figure nascent polypeptides will be immunoprecipitated upon partition the translational complexes in SPERT. One partition ribosome complexes or mRNA.polypeptide

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useful for the development of polypeptide ligands which has an immuno-dominant domain for which polyclonal sera domain (the "handle"). This modification is especially segment of a fusion protein in which the amino terminus binding antibodies which recognize the immunoreactive galactosidase, for example) and the carboxyl-terminal is available. Where immunoprecipitation is employed, portion is the protein of interest. It will also be recognize immunoresistant domains of a protein which useful for developing polypeptide ligands against a complexes or mRNA polypeptide copolymers that react directly with the antibodies, prior to selection. it will be advantageous to discard any ribosome is the fragment of a common protein (beta-

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mixture or test mRNA.polypeptide copolymers mixture and target antibody will be retained on the column as part separating polypeptides bound to targets, particularly of the ribosome complex or mRNA polypeptide copolymer mRNA's that encode a polypeptide that binds to the affinity column which retains the immune reactive proteins, are available to the art. For example, passing the immune complexes through a protein A and unbound coding mRNA's can be washed from the immunoprecipitation of the test ribosome complex polypeptide-containing complexes as the column. Alternative partitioning protocols for binding and partitioning can be achieved by

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column.

location for randomization and SPERT-based isolation of or multiple strands, and either structure can be formed loops or hairpins, are related to RNA hairpin loops and cysteines represent another means by which to construct those structures include alpha helices and beta sheets Interestingly, protein loops may be a powerful novel ligands. When inspecting protein structures in on the loop primary sequences and contacts with other the loops are set by the secondary structures but the exact loop structures are idiosyncratic and dependent connectors between such secondary structures, called RNA pseudoknots in that the locations of the ends of randomized and put through SPERT should provide vast loops; similarly, zinc fingers and copper or other detail, only secondary structures are predictable; metal "fists" also provide other kinds of loops. elements of the protein. Loop sequences, when structural libraries. Disulfide bonds between with parallel or anti-parallel peptides. The

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Effective partitioning can be carried out with pure or impure target preparations. In cases where target preparations. In cases where target preparations are impure, selectivity can be enhanced by strategies that enhance the binding of ligands to the desired target, or which specifically elute desired ligands or prevent their binding. The latter approach is subtractive. A known ligand can block binding of any polypeptide that can bind the target so that the desired polypeptide is partitioned by elution and unwanted polypeptides are retained on the column.

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Optionally, chemical or enzymic modifications of the polypeptide can be introduced posttranslationally. The process for making such modifications should not disrupt the ribosome complexes or mRNA.polypeptide.copolymers. An important type of post-translational modification is oxidation to form

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disulfides in sequences that contain two or more cysteines. Particularly for small polypeptides, disulfide bonds are especially advantageous to lock in a desired conformational state so that a rigid structure having high specificity and binding affinity for a target can be achieved. (See, e.g., Olivera, B. M., et al. (1990) Science 249:257-263.

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functions, i.e., capable of being functionally "off" or covalently influence tertiary structure of the nascent selecting polypeptides having conditionally reversible nucleotides, flavine nucleotides, porphyrins, thiamine degrade the coding mRNA or nascent polypeptide should be avoided. A modifier can be included in the buffer potential binding activities of any candidate mixture need to maintain stability of the ribosome complexes. polypeptide can exhibit different binding properties or medium during partitioning. Alternatively, SPERT modifiers during partitioning is only limited by the polypeptide. In particular, metal ions such as Ca biological functions, can interact with the nascent Modifiers which disrupt ribosome complexes or which modifications include introducing factors that non-Other forms of post-translational structure coordination complexes with amino acid side chains. Mg⁺, Mn⁺, Zn⁺, Fe⁺⁺, Fe⁺⁺, Cu⁺⁺ and Mo⁶⁺ can affect "on", depending on the presence or absence of the Similarly organic compounds such as nicotinamide than an unmodified polypeptide. The use of such modifier. Configurational modifiers need not be naturally-occurring compounds. The use of such configurational modifiers enhances the range of of polypeptides. Also, it affords a means for phosphates, serotonin, and the like, including polypeptide to modify its 3-dimensional folded inhibitors, agonists and antagonists of known configuration. As thus modified, the nascent polypeptide folding configuration by forming 10 20 12 25 30 35

itself can be used to pre-select polypeptides which bind the modifier as a target after which the candidate mixture of selected modifier-binding polypeptides can be further selected, via SPERT, for binding the

ultimate target.

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Sequence variation in the test coding mRNA mixture can be achieved or increased by mutation. For example, a procedure has been described for efficiently mutagenizing nucleic acid sequences during PCR amplification (Leung et al. 1989). This method or functionally equivalent methods can optionally be combined with amplification procedures in the present invention.

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Alternatively, conventional methods of DNA mutagenesis can be incorporated into the nucleic acid amplification procedure. Applicable mutagenesis procedures include, among others, chemically induced mutagenesis and oligonucleotide site-directed mutagenesis.

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approach is to replace the variable (or hypervariable) partitioning conditions, as will be apparent to those sequence in the mRNA that encodes the protein. Since modifications, alterations and improvements on known sequences synthesized <u>de novo</u>. In particular, SPERT The starting mRNA mixture is not limited to sequences conserved and others varied, the logical chance of altering function. The proper choice of many known proteins belong to families having some regions with randomized sequence, to maximize the replaced by a corresponding segment of randomized skilled in the art, results in selection for the proteins. A segment of the natural sequence is can be used to modify the function of existing desired functional variant. In this way, proteins can be achieved.

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To proceed to the amplification step when utilizing ribosome complexes, coding nucleic acids must

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after partitioning. This process must be done without complex-target pairs by phenol extraction or by phenol such that the mRNA is eluted. The coding mRNA can be subsequent amplification is selective for the mRNA's embodiment, selected coding RNA molecules are eluted other eluant capable of disrupting the ligand-target be released from the target-bound ribosome complexes combined with a protein denaturing agent such as 7M because the primers used for cDNA synthesis and PCR result in amplifiable nucleic acids. In a specific from a column using a high ionic strength buffer or bond. Alternatively, the ribosome can be denatured amplification are complementary only to a conserved chemical degradation of the coding mRNA's and must removed from ribosome complexes or from ribosome urea. Although ribosomal RNA is also extracted, seguence in the mRNA's and not to ribosomal RNA. ហ 15 2

As the translation of randomized mRNNs proceeds during the SPERT protocol, the growing polypeptide makes its way from the peptidyl transferase site within the large ribosome subunit toward the cytoplasmic solvent. The peptidyl transferase site is an intrinsic activity of the large ribosome subunit from all organisms; that site has been defined functionally but its precise location within the ribosome is unknown. However, the distance between that site and the cytoplasmic solvent also is known to be about 30 to 40 amino acids in length.

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portion of the nascent polypeptide (whose properties are selected during the procedure) should be "outside" the ribosome in order for partitioning of the ribosome complex to fully utilize the properties of the randomized polypeptide. A C-terminal trailer sequence is preferably incorporated into the translated polypeptide to insure that the randomized sequence is fully exposed after translation. From the work of

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if the amino-terminus of a growing polypeptide contains a hydrophobic domain of about 20 amino acid residues, a nascent polypeptide of about 50 residues has been shown those preferred embodiments of SPERT utilizing ribosome by randomized mRNA that is about 30-40 codons (that is, SPERT, itself, can be used to determine optimum trailer Smith et al, (PNAS, 75:5922, 1978) and Malkin and Rich (J. Mol. Biol., 26:329, 1967) for both prokaryotes and eukaryotes: about 30 to 40 amino acid residues remain see Kurzchalia et al, Nature 320:634, 1986). Thus, in complexes, the randomized polypeptide will be encoded about 90-120 nucleotides) upstream from the codons at within the ribosome during translation. Furthermore, interact with a membrane by hydrophobic interactions, length for a given partitioning system. The sequence polypeptide purification, as a reporter activity for trailer sequences can be used effectively, and that function, such as more stability in the translation understood that both longer and shorter C-terminal complex, ease of in vitro manipulation, subsequent which the translation complex stalls. It will be of mRNA and encoded polypeptide in the C-terminal trailer can be designed to have any other desired to be enough to allow the translation complex to diagnostics, cell entry, etc.

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Polypeptides selected by SPERT can be produced by any peptide synthetic method desired. Chemical synthesis can be accomplished since the amino acid sequence of the polypeptide is readily obtainable from the nucleotide sequence of the coding mRNA. Since cDNA from the coding mRNA is available, the polypeptide can also be made by expressing the cDNA in a suitable host cell.

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SPERT offers, as noted above, an opportunity to sample as many as 10^{18} peptide sequences during a rigorous experiment with a particular target. As such SPERT may be compared with $\underline{1}$ \underline{v} \underline{v} technologies aimed

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phage display systems depend, in their present form, on a transformation step with either plasmid or phage DNA, for more than five years (see, Smith, Science <u>228</u>:1315, e.g., Charbit et al., EMBO J. 5:3029, 1986; Parmley et properties. These technologies, lumped together under al., Gene 73:305, 1988; Scott et al., Science 249:386, with bigger volumes and more difficulties. SPERT thus 1990; Devlin et al., Science 249:404, 1990; Cwirla et peptides to be searched easily, and perhaps 10^{11} or so the name "phage display systems", have been available al., Proc. Natl. Acad. Sci. 87:6378, 1990). Because the intrinsic depth of those systems is less than in 1985) and widely appreciated in the last year (See, SPERT. Phage display systems allow 10^9 different has a value for looking rigorously through large at uncovering peptides with specific binding libraries.

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display systems have a disadvantage in common, at least and which is extremely large relative to the peptide of formally. In SPERT the peptide of interest is held by the ribosome, a machine that contains its own proteins interest. Similarly, in the phage display systems the Both SPERT, as defined thus far, and the phage improved system would provide the peptide of interest bound to an encoding nucleic acid (to achieve reverse systems will yield a peptide of interest with careful process. The large entities also severely limit the translation) free of any other large, proteinaceous peptide of interest protrudes from a phage particle which is also relatively extremely large and which contains its own proteins. Although each of these particle and the ribosome add limitations to these systems other than in the partitioning step of the number of random peptides that may be practically partitioning of the bound peptide from all other peptides bound to ribosomes or phage capsids, an components. As described above, the large phage

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generated and tested in the screening process.

peptides becomes covalently or very tightly attached to an alternate embodiment, this invention contemplates a one end or the other of the mRNA that encodes it to SPERT lends itself to such an improvement. simple and general mechanism by which a non-random portion of each peptide within the collection of form mRNA polypeptide copolymers.

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allows the ribosomes in the translation mixture to have to lower RNase levels. Alternately, various techniques 3) the reaction between the nascent polypeptide and the problem may also be alleviated by using mutant strains a high turnover can be useful. The in vitro reactions familiar to those skilled in the art are available for 2) additional reagents should be relatively small; and mRNA.polypeptide copolymers include the following: 1) the mRNA must either occur before the ribosome complex interaction over dissociation of the proximal species; should be as free as possible from RNases. The RNAse the interactions between the nascent polypeptide and specific systems that could be employed to generate mRNA should be relatively efficient (i.e., at least mRNA.polypeptide copolymers. Any such system that is disrupted, or at a rate that highly favors the making the mRNA nuclease resistant. Additional There are an almost unlimited number of criteria for effective systems for forming about 5% or greater).

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Activation of the 5' end of the mRNA species and the Norganic chemical type reactions between the species; 3) terminus of the peptides to promote relatively simple onset of translation; and 4) tRNA crosslinking of the employed to generate mRNA.polypeptide copolymers will A nonlimiting catalog of methods that can be Adapted post-translational modification systems; 2) Attachment of the peptide to the mRNA prior to the generally fall into the following categories: 1)

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nascent polypeptide and the mRNA. Various embodiments systems would also be obvious to those skilled in the of each of these systems is described below. The design of additional embodiments of these general

Post Translational Modification Systems

art.

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harming the RNA for other functions. Thus each mRNA in one of a number of small reagents affixed to the 5' end ribosomes are indifferent to the chemical nature of the nucleoside triphosphates are the source of all internal attached to the 5' end without difficulty, and without Burgin et al., EMBO J. $\underline{1}$:4111, 1990, for example), the In one embodiment the collection of mRNAs used In SPERT is synthesized using T7 RNA polymerase and 5' the collection could have, for example, biotin or any guanosine phosphono monothioate for initiation (see, of the RNA. Alternatively, mononucleotides labeled with biotin could be used to initiate transcription. residues during transcription. Organic tags may be upstream of the initiating AUG and as long as those nucleotides contain appropriate sequences to cause The 5' end of the RNA would certainly not preclude 5' end as long as enough nucleotides are present translation by bacterial ribosomes, since those monothioate is incorporated only at the 5' end; initiation to occur. 12 20 25 ដ

the 5' end of each mRNA. Known peptide sequences (such covalently bound to the chemical adduct positioned at as avidin) might be used if biotin were the chosen 5' make covalent the interaction between the peptide and downstream from the AUG, also fixed, encode a peptide tag. In one example, a biotin ligase may be used to (Cell, 58:427, 1989); Reed and Cronan (J. Bio. Chem., the biotin at the 5' end of the mRNA. See, Cronan, that has an extremely high affinity for or can be According to this embodiment, the codons

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Many suitable pairs of chemical adducts and fixed peptide sequences have been identified, and are known to those skilled in the art. For example, certain polypeptides contain lipoylation sites, and the post-translation modification would utilize the lipoylation system. See, Rucker et al., (FASEB J., 2:2252-61, 1988); Ali et al., (Mol. Microbiol. 4:943-50, 1990). For other post-translational modification systems, see, PCT Patent Application PCT/US90/02852 (published November 29, 1990, WO 90/14431).

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ribosome, the most likely 5' adduct to be bound by that that the distance between the nascent, emergent peptide fixed. The length of the 5' end of each mRNA (that is, site are needed to enhance the binding reaction in <u>cis</u>) and the 5' end of the mRNA of another can be determined methionine). Again, with respect to biotin and biotin collisions between the nascent peptide of one ribosome easily without undue experimentation. This last point this case, randomized peptide sequences downstream of how many nucleotides upstream of the ribosome binding about 200 angstroms in diameter, so it may be assumed (from the large ribosome subunit) and the emergent 5' will never be more than 500 angstroms apart and could nascent peptide with respect to its own 5' adduct in adduct of the mRNA (from the small ribosome subunit) encoding that exact peptide (which will include, in peptide sequence will be the 5' adduct on the mRNA is clear from a simple calculation. Ribosomes are be much less. The calculated concentration of the Since the ribosome concentration in many cell-free ligase, the first collisions will be irreversibly scenario, and could be more than 100 times higher. <u>cis</u> is higher than 3 micromolar for a worst case As the nascent peptide emerges from the and the concentration of ribosomes that allow the fixed peptide adjacent to the initiating

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translation experiments is sub-micromolar, it is not difficult to preclude scrambled binding between nascent peptides and 5' mRNA adducts on other ribosomes.

are used for the linkage). More complete purifications mRNA.polypeptide copolymer with antibodies against that of the copolymer prior to partitioning with target are disassociate the ribosomal subunits. ATA, poly U, or merely effective copolymers when very high affinities eliminate the ribosomes and many of the proteins from peptides that partition with a target, the cell-free obvious. For example, hybridization to column-bound copolymer (such copolymers may be truly covalent or As translation ends, after mRNA polypeptide epitope would do, thus allowing purification of the reaction may be treated with puromycin and EDTA to other non-amplifiable RNAs may be added to prevent fractionation may then be used to enrich for small additional sequence for this purification; a small material, and/or high speed centrifugation would copolymer formation and prior to enrichment for subsequent elution would give full purification. the cell-free system from the mRNA.polypeptide complementary DNA (to one end of the mRNA) and Similarly, the fixed peptide could include an rebinding of mRNAs to the ribosomes. Size

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The mRNA-polypeptide copolymer is partitioned as in the ribosome complex examples, and the bound mRNA amplified via cDNA synthesis and PCR, as always extending the cDNA to create again the T7 promoter sequence for the next round of SPERT. The peptide attached to the 5' end of the mRNA may cause the 3' end of the cDNA to be a bit shorter than in the absence of peptide, but PCR easily accomplishes the full restructuring of the DNA for subsequent transcription, in this case initiated once again by phosphono monothioate nucleotide for adding the small organic

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molecule needed for linkage.

In this alternate embodiment of SPERT, the peptide is directly linked to the encoding nucleic acid and is partitioned to target (or reacted in any other way described for SPERT) with only the encoding nucleic acid available (along with the peptide collection) for that target. The very large ribosome or phage capsid no longer obscures the partitioning reaction in any way.

Activation of 5' end of mRNA and N-terminus of peptide.

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The post-translational modification systems described above generally require an enzyme to facilitate the reaction between the nascent peptide and the mRNA. According to this embodiment, the modifying enzyme is eliminated, and relatively simple chemical reactions are relied on to form the copolymers.

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In one embodiment of this system, sulfur-halide chemistry is employed. Sulfur may be incorporated on the 5' end of the mRNA using the T7 RNA polymerase and monothiate for initiation as described above. A halide can be incorporated on the N-terminus of the peptide by use of N-haloacetyl-met-tRNA^{fmet} (Pellegrini <u>et al.</u>, (Proc. Natl. Acad. Sci. USA, 69:83741, 1972); Sopari, <u>et al.</u>, (Biochemistry, 13:5432-39, 1976)). This combination would result in spontaneous nucleophilic substitution to form a thioether linkage between the nascent polypeptide and the mRNA. In order to avoid reaction of the halo-acetyl group with DTT in the translation mixture, or with cysteine residues in ribosomal proteins, it is preferred that the chloro acetyl functionality be utilized.

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In a further embodiment of this process, it may be desirable to accelerate the reaction between the nascent polypeptide and the mRNA by introducing a "chaperone" RNA sequence. The chaperone acts as a catalyst to facilitate the nucleophilic substitution

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reaction. An appropriate chaperone sequence may be easily selected by one skilled in the art utilizing the SELEX technology. A useful chaperone may be selected by placing a stretch of random noncoding RNA adjacent the 5' GMPS mRNA, and collecting those sequences capable of reacting with the halo-acetyl N-terminal polypeptide. This reaction could be further facilitated by selecting fixed amino acids at the N-terminal end that would present a probable nucleic acid interaction site. In further embodiments, the chaperone could be an RNA or protein acting as a true catalyst to facilitate the reaction.

Pre-coupling of mRNA to Peptide.

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In one embodiment of the formation of mRNA.polypeptide copolymers, the mRNA may be coupled to the nascent polypeptide before translation is initiated. In one embodiment, this pre-translational coupling would occur by attaching the 5' end of the mRNA to the X-amino group of methionine on met-tRNA via a covalent linker. As translation proceeds, the initiating methionine is already attached to the mRNA at the initial amino acid sequence.

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25 tRNA Crosslinking of Message and Peptide.

linkage is created between peptidyl-tRNA and mRNA. A specific embodiment of this system is based on studies of the photoreaction between the "Y" base of yeast tRNA^{phe} and mRNA. See, Matzke et al., (Proc. Natl. Acad. Sci. USA, 77:5110-14, 1980). See also, Steiner et al., (Nucl. Acids. Res. 12:8181-91, 1984) (demonstration that tRNA can undergo peptidyl transfer and translocate normally from A-site to P-site after being crosslinked to mRNA); Paszyc et al. (Nucl. Acid. Reg. 6:385-97, 1979). A nonsense suppressor containing the Y base may be used that will crosslink to the

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Continuous irradiation of this system during translation would yield photocrosslinked mRNA-polypeptide copolymers. An advantage of this embodiment is that there would not be any constraints on the peptide or message.

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It is an important and unexpected aspect of the present invention that the methods described herein can be employed to identify, isolate or produce polypeptide molecules which will bind specifically to any desired target molecule. Thus, the present methods can be employed to produce polypeptides specific for binding to a particular target.

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Proteins contain within their primary sequence the information required to form an extraordinary variety of three dimensional shapes as is well known in the art. From this variety of potential shapes, along with the charge and/or hydrophobic qualities of amino acids, comes the potential for protein functions that are used in the biosphere. Proteins provide catalysis when embodied as enzymes; proteins can provide stable biological structures, for example, when used to construct spores, membranes, or viruses; and proteins can provide binding to a variety of targets, with appropriate affinities and kinetic parameters to allow life.

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Nevertheless, this vast potential in chemical activities, including the extreme potential inherent in the mammalian immune system, has actually been explored rather narrowly by organisms. This fact can be noted with a simple calculation. If the average length of a protein is 300 amino acids, and if there are twenty

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present invention provides the means to explore protein the number of possible sequences of proteins of average history and the relatively short age of the earth. The established by the number of particles in the universe. The tiny fraction of so-called sequence space that has natural amino acids used to construct modern proteins, particles in the universe are in the range 10^{80} , while estimates for the number of proteins ever explored in the entire history of the earth are in the range 10^{10} largely of individual entities that have never before ligand can be learned from the nucleotide sequence of its encoding mRNA, making tedious amino acid sequence been explored by biology is a result of evolutionary limitations, while continuing to respect limitations isolate polypeptide ligands with any desired quality sequence space without historical and evolutionary size is 20^{300} or ~ 10^{400} . Estimates of the number of from vast mixtures of protein sequences comprised existed. The amino acid sequence of the selected The invention provides the means to identify and determination unnecessary.

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proteins or peptides having similar function. In most instances, SPERT-selected polypeptides will be smaller leader of about 10 amino acids, for a total length of than naturally-occurring proteins typically having a size of from 4-100 amino acids, preferably from 4-50 amino acids selected from randomized sequence of the same length, and also having a C-terminal trailer of about 30-40 amino acids and, optionally a N-terminal Even where the binding functions selected by enzymes and all antibodies, for comparison, IgG has selected by SPERT will resemble naturally-occurring about 100 amino acids, corresponding to a molecular molecular weight of about 150kd. Furthermore, many there is no reason to expect that the polypeptides SPERT have known naturally occurring counterparts, weight of about 11kd. This is smaller than most

polypeptide ligands of the invention will function when space, it is expected that most polypeptide ligands of occurring, and typically differ in amino acid sequence freed by N- and C- terminal trailers. Therefore, the Additionally, given the vastness of sequence randomized coding is designated the "binding segment" and molecular size from naturally-occurring proteins. That portion of the amino acid sequence arising from natural proteins, and preferably less than 30% amino conveniently ranging from about 4-100 amino acids in the invention will have less than 50% homology with final product can be as small as 4-50 amino acids. length, preferably from about 15-50 amino acids in herein. The binding segment can be of any length, polypeptides of the invention are non-naturallyacid homology with natural proteins.

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than antibodies and they can be generated by techniques antibodies, they can be selected to bind at any desired sites not recognized by antibodies, they can be larger live animals or cell culture techniques. Applications invention have significant advantages over antibodies: similar to those of antibodies can be isolated by the application. However, the polypeptide ligands of the epitope or combination of epitomes, including binding methods of the present invention. Such polypeptides including higher affinities than are obtainable with that operate entirely in vitro, without the need for or smaller and have different solubility properties A polypeptide ligand of the invention in a number of ways functionally resembles an antibody. molecules from any source; purification of target Polypeptide ligands which have binding functions qualitative or quantitative detection of target polyclonal or monoclonal antibodies have found they can be selected for any desired affinity, are generally useful in applications in which of polypeptide ligands include the specific,

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to function via binding to nucleic acid sequences, such nucleic acid sequences which are known to bind proteins The methods of the present invention related to the use that bind targets for which other proteinaceous ligands are known. For example, a number of proteins are known proteins. The methods of the present invention related molecules are preferably proteins, but can also include sites, for example, has been employed in the detection, polypeptide; and various therapeutic methods which rely of polypeptide ligands can generate novel polypeptides nucleic acid binding proteins to bind to their natural novel nucleic acid binding ligands having affinity for proteins. Novel, non-naturally-occurring polypeptides cell surfaces or viruses, through specific interaction to the use of polypeptide ligands can be used to make which bind to the same binding sites of nucleic acids also be employed to affect the function, (for example with conventional antibodies, polypeptide ligands can production, since SPERT is a wholly in vitro process. be employed to target biological structures, such as therapeutic agent to a specific target site. Target peptidoglycans and a variety of small molecules. As tolerance, as are conventional antibodies. Also, as below, certain polypeptides isolatable by SPERT can can be developed using SPERT. As will be discussed noted, polypeptide ligands of the invention do not require animals or cell cultures for synthesis or as regulatory proteins which bind to nucleic acid operator sequences. The known ability of certain advantageous in that they are not limited by self quantitation, isolation and purification of such molecule's based on their specific binding to the with a molecule that is an integral part of that and to nucleic acid sequences not known to bind biological structure. Polypeptide ligands are on the specific direction of a toxin or other among others carbohydrates, nucleic acids, 32 S2 30 20 ß 10 12

inhibit, enhance or activate) specific target molecules or structures. Specifically, polypeptide ligands can be employed to inhibit, enhance or activate the function of proteins and of nucleic acids.

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Polypeptide ligands of small molecules are particularly molecules which will bind specifically to a particular change in the target. The range of possible catalytic It is a second important aspect of the present also include, among others, carbohydrates and various small molecules to which specific polypeptide binding interacting with their natural ligands. For example, polypeptide ligand that binds the enzyme's substrate. function of a small molecule, by catalyzing a chemical employed to identify, isolate or produce polypeptide quantities of normal metabolites can be detected and activities is at least as broad as that displayed by controlled substances, bound metabolites or abnormal molecule. In this aspect, the target molecules are again preferably proteins or nucleic acids, but can measured using polypeptide ligands of the invention. invention that the methods described herein can be can be achieved. Polypeptide ligands that bind to quantitative assays. For example, the presence of precipitate or bind ligand-target pairs to a solid phase matrix in a diagnostic assay. A polypeptide useful as reagents for diagnostic tests, or other Antibodies to polypeptide ligands can be used to target molecule and affect the function of that ligand having catalytic activity can affect the the activity of an enzyme can be affected by a sequestering them or by preventing them from small molecules can affect their function by natural proteins.

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The strategy of selecting a ligand for a transition state analog of a desired reaction is one method by which catalytic polypeptide ligands can be selected. Polypeptide ligands with high affinity for

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transition-state analogues are likely to have enzymatic [Pollack et al., Science 234:1570 (1986); Tramantano et 237:1041 (1987); Janda et al., Science 241:1188 (1988); catalytic activities, including acyl-transfer reactions 110:5593 (1988)], carbon-carbon bond cleaving reactions antibodies directed against transition-state analogues. Schultz, P.G., Science 240:426 (1988); Benkovic et al., peptide cleavage [Iverson and Lerner, Science 243:1184 sequences and structures that can be explored by SPERT Proc. Natl. Acad. Sci. 85:5355 (1988)], carbon-carbon 110:4841 (1988); Hilvert and Nared, J. Am. Chem. Soc. [Cochran et al., J. Am. Chem. Soc. 110:7888 (1988)], Science 244:437 (1989)]. The number of polypeptide al., Science 234:1570 (1986); Jacobs et al., J. Am. Chem. Soc. 109:2174 (1987); Napper et al., Science bond formation [Jackson et al., J. Am. Chem. Soc. activity, as has been demonstrated for monoclonal (1989)], and ester bond hydrolysis [Janda et al., far exceed those available in the immune system. These antibodies have exhibited a wide range of

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Enzymes are evolved using SPERT and starting randomized sequences corresponding to about 50 amino acids, as in Example 3. Enzymatic polypeptide ligands of small size are entirely unanticipated by the present understanding of enzymology; enzymes are always much larger in nature than the scientist expects. The specific transition state analogues used are drawn from the literature cited above. Among the reactions probed by the monoclonal antibody-enzymes are some which lead to the breakdown of toxic waste products, including chemicals with chlorine-carbon bonds and carbon-carbon bonds in ring structures like those found in benzene and polychlorinated phenols.

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The binding selection methods of the present invention can be combined with secondary selection or screening to identify ligands capable of modifying target molecule function upon binding. The large

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to isolate or produce polypeptide ligands which bind to and modify the function of any protein or nucleic acid. protein. The methods described herein can be employed protein, i.e., disrupt transport of small molecules or populatión of variant amino acid seguences that can be polypeptide sequences can be found that have a desired invention are useful for selecting polypeptide ligands binding capability and that function to modify target binding to receptors; affect the formation of protein protein subunits; and modify transport properties of which can selectively affect function of any target catalytic activity of target enzymes, i.e., inhibit invention can be employed to identify, isolate or catalysis or modify substrate binding, affect the functionality of protein receptors, i.e., inhibit binding to receptors or modify the specificity of It is contemplated that the method of the present multimers, i.e., disrupt quaternary structure of produce polypeptide molecules which will affect molecule activity. The methods of the present tested by SPERT enhances the probability that ions by proteins.

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structure, etc. Those of ordinary skill in the art are selection or screening methods that are compatible with combined with SPERT include among others selections or screens for enzyme inhibition, alteration of substrate able to select among various alternatives those Secondary selection methods that can be binding, loss of functionality, disruption of the methods described herein.

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employs a molecule known, or selected, for binding to a polypeptides which bind to a particular functional or An embodiment of the present invention, which desired site within the target protein to direct the is particularly useful for identifying or isolating active site in a protein, or other target molecule, selection/amplification process to a subset of

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sequence encoding the known bridge-binding oligopeptide the selection/amplification steps repeated to select an even longer binding sequence. Iteration of these steps longer binding sequence, which is anticipated to either longer binding sequence can then be introduced near the is incorporated near the randomized region of the test amino acid. In this embodiment a molecule which binds a polypeptide sequence known to bind to a desired site randomized region of all polypeptides being tested for covalently linked to the "anchor" molecule that binds select for those polypeptide sequences that bind most site within the target molecule. In a simple example, binding. SPERT is then used to select those variants, specifically to the target can thus be selected. The randomized region of the polypeptide test mixture and within a target molecule. Another embodiment of such an iterative "walking" procedure, employs an "anchor" to the target also binds to the target molecule. The procedure allows the selection of polypeptides highly all of which will contain the known binding sequence, oligopeptide of known sequence. The bridge molecule inhibitor of a target enzyme, is chemically modified polypeptide ligands that bind at or near the desired which bind most strongly to the target molecule. A (i.e., incorporation of selected sequence into test molecule which is not necessarily a polypeptide or improved or more specific binding) can be repeated specificity is achieved. This iterative "walking" specific for a particular target molecule or site molecule which in turn is known to be bound to an nucleic acid mixture. SPERT is then performed to bing more strongly to the target molecule or more such that it can be covalently linked to a bridge mixtures followed by selection/amplification for to a desired target, for example a substrate or in a target molecule is incorporated near the until a desired level of binding strength or 30 32

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The iterative walking procedure can then be employed to isolation of polypeptide ligands that bind at or near a strongly to the target molecule/bridge/anchor complex. it is expected that the "anchor" method in combination desired site within a target molecule. In particular, specificity of binding to the target. The use of the polypeptides which are highly specific inhibitors of "anchor" procedure is expected to allow more rapid with iterative "walking" procedures will result in select or produce longer and longer polypeptide molecules with enhanced strength of binding or protein function.

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or mRNA polypeptide copolymer is, in principle, capable of binding to target molecules and of being partitioned differentiated from polypeptide binding once the ligand has been selected and both the selected polypeptide and concurrently with nascent polypeptides. In particular, binding studies where the two are not part of the same ribosome complex. Comparative studies of the relative For a given target it can not be predicted whether RNA or peptide will give more useful ligand solutions, and 07/714,131, the translated mRNA of a ribosome complex protein targets will yield a tight-binding RNA ligand. function that currently exists in living cells. This SPERT cycles may prove to be surprisingly robust. As described in the SELEX applications, large numbers of In accordance with the teachings of copending thus SPERT can be seen as an improvement to the SELEX direct comparison between RNA and peptide during the its coding mRNA are available for independent direct rather than a polypeptide. Binding of mRNA can be chromatography, the selected ligand can be an RNA, applications Serial No. 07/536,428 and Serial No. importance to understanding the specialization of frequency of RNA ligands and polypeptide ligands selected by SPERT are of fundamental biological where partitioning is accomplished by affinity

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conserved RNA sequence or structure is found, while the peptide solutions will force the RNA solutions to have will be indifferent to the reading frame in which the immediately. For example, the RNA ligand solutions application because when RNA yields the best ligand solutions the data will lead to that conclusion a common sequence in the same reading frame.

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of the buffer medium used during partitioning. As will membrane affinity can be included as a property, either For example, during partitioning, stability to certain certain protease is desired, that protease can be part avoided. Other desired properties can be incorporated hydrophobicity, or by biasing the randomized coding to If a polypeptide which is stable in the presence of a conditions which disrupt ribosome complexes should be end product can be included as a selection criterion. selected for other properties in addition to binding. conditions of the desired working environment of the understood by those skilled in the art. For example, by employing a N- or C-terminal trailer having high favor the amino acids with lipophilic side chains. directly into the polypeptide sequence as will be be understood, when utilizing ribosome complexes The polypeptides of the invention can be

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resistent to a virus infection, for example, by causing nucleic acid or a key viral protein. In principle, any in <u>vivo</u> synthesis of a polypeptide ligand of the viral The coding nucleic acid concomitantly selected by partitioning nascent polypeptides as described, is functionality contributed by a polypeptide ligand of polypeptide. A transgenic organism can be rendered organisms. The transformed organism is then useful useful in its own right to transform host cells or for, e.g., fermentation production of the selected organism. Methods known in the art can be used to the invention can be bestowed on a suitable host combine the coding region with a promoter,

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polyadenylation signal functional in the intended host, followed by incorporation into a suitable vector for transformation, all as known and understood in the art.

EXAMPLES

The techniques and methods used in the ensuing examples are published and known in the art. Together with adaptations and modifications known to those of ordinary skill in the art, the procedures not specifically referenced herein are available from known reference works. In addition to Sambrook et al., (1989) <u>Supra, Genetic Engineering</u>, Plenum Press, New York (1979); Weir, (ed.) (1986) <u>Handbook of Experimental Immunology in Four Volumes</u>, 4th Ed, Blackwell Scientific Publications, Oxford; and the multivolume <u>Methods in Engymology</u> published by Academic Press, New York. Polymerase chain reaction techniques are described in <u>PCR Protocols</u> (Michael A. Innis, et al. eds.) (1990) Academic Press, Inc.

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Throughout examples 1-9, reference is made to Tables 1 and 2. Table 1 lists oligonucleotide sequences used for preparing mRNA candidates. Table 2 lists the same sequences together with explanatory notes showing functional domains. Sequences in capitals are chemically synthesized, sequences in lower case letters are complementary sequences made enzymatically by DNA polymerase. The Examples could be adapted by those of ordinary skill in the art to generate mRNA-polypeptide copolymers as taught herein without undue experimentation.

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Example 1. Direct Immunoprecipitation of Ribosome Complexes: Polypeptide Ligands Directed Toward Immunoglobulin Molecules.

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The method of the invention is used to select novel polypeptides that bind the antibody of an epitope

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mice which are the fl progeny of a cross of NZB and NZW into a restriction site to provide an mRNA encoding the promoter sequence and a ribosome binding site which is and 2 and in Figure 8. A 3' fixed sequence is placed lacking stop codons (for ca. 30-35 amino acids) shown histone H2B protein. To make mRNA encoding candidate site is synthesized and cloned using oligonucleotides addition, as shown in Figure 1, a 3' primer annealing site (sequence 3) is provided so that cDNA synthesis ribosomes, terminating in a restriction endonuclease having the sequences shown as sequence 1 in Tables 1 commonly recognized by the antisera from autoimmune contiguous amino acids at the amino terminus of the C-terminal trailer sequence of ca. 100 nucleotides polypeptides, a 5' fixed sequence composed of a T7 as sequence 3 in Tables 1 and 2 and Figure 8. In parents (Portanova et al., J. Immunol. 144, 4633 (1990). The known epitope consists of about 10 can be accomplished on the mRNA recovered from recognized by both prokaryotic and eukaryotic partitioned ribosome complexes. Ŋ ដ 12 20

Synthesis of randomized oligonucleotides is carried out the reaction mixtures contain, on a mole percent basis, sequence of 45 nucleotides (corresponding to 15 codons) bounded by restriction endonuclease recognition sites, with a reactant mixture for each nucleotide position. bounded by specific sequences that include those two The randomized polypeptide insertion site is using an Applied Biosystems DNA synthesizer provided To partially compensate for the amino acid sequence bias inherent in the redundancy of the genetic code, First position, C-20%, T, A, and G-30% each; Second in this example EcoRI and PstI. A single-stranded the following composition of bases for each codon: oligonucleotide is synthesized with a randomized position, C-15%, A-35%, T and G-25% each; Third restriction endonuclease sites (Sequence 4a).

position, T, C, A and G-25% each. Using a nucleic acid primer that is complementary to the fixed 3' end of the randomized oligonucleotide, randomized double-stranded DNA is created with the action of DNA polymerase. The products are digested with the two restriction endonucleases and ligated between the 5' fixed sequence and the 3' fixed sequence discussed above. In vitro transcription of these ligated templates using T7 RNA polymerase (Bethesda Research Laboratories,

Gaithersburg, MD) provides mRNA templates for in vitro translation. A rabbit reticulocyte lysate system (BRL) is used to translate the mRNA templates in vitro, using standard reaction conditions. Such translation of these transcripts results in a variety of ribosomal complexes (mRNA-nascent polypeptide-tRNA-ribosomes) that are identical except for the randomized region of the nascent polypeptide.

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Antibodies (IgGs), Portamova et al., Subra, which recognize the H2B histone epitope are added to the in vitro translation mixture. Immunoprecipitation of the immunoreactive ribosome complexes partitions the mRNAs species that encode the highest-affinity polypeptide ligands in the population (see Figures 3 and 4). Immunoprecipitated complexes are separated by low speed centrifugation. cDNA is synthesized from these mRNAs and is used via PCR to provide template for further cycles of transcription, translation, immunoselection and cDNA synthesis.

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Clones are isolated as described in Application 07/536,428, June 11, 1990, incorporated herein by reference. The individual polypeptide products are over- produced and purified and tested, using standard techniques for reactivity to the anti-H2B histone antibodies. In addition, the polypeptide ligands are challenged competitively with authentic histone H2B-derived epitomes to discover which polypeptide ligands bind to the same portion of the antibodies as the true

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epitope. Among the polypeptides isolated that bind the antibody are found those having less than 50% sequence homology with the H2B histone epitope. Other antibody binding sequences are identified having less than 30% homology with the H2B histone epitope. Other polypeptide ligands of the antibody do not compete for the H2B epitope binding site.

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Example 2. Diagnostics using the polypeptide ligands of Example 1: An assay for anti-H2B antibodies in the progeny of NZB X NZW

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Auto-immune diseases result from the elaboration of an inappropriate antibody molecule with reactivity toward a normal cellular component (often a protein, but sometimes a nucleic acid, as in Systemic Lupus Erythematosis - SLE). Polypeptide ligands generated through the SPERT protocols in Example 1 are aimed at diagnosis of mouse "Lupus" in the offspring of NZB X NZW mice. SPERT is used to identify and obtain a reagent ligand for the diagnostic recognition of the auto-antibody that recognizes the histone H2B epitope.

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As in Example 1, ribosome complexes are treated with the auto-antibody to partition reactive polypeptides from non-reactive polypeptides resident (as nascent polypeptides) in ribosome complexes. The auto-antibodies are used to precipitate the ribosome complexes containing polypeptides that fit into the active site of the antibody. The most avidly bound polypeptide emerges from repeated SPERT cycles.

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The most avidly bound polypeptide ligand does not resemble in detail the epitope identified as the portion of the target that reacts with the antibody.

Auto-immune diseases are triggered by unknown antigens, which are not necessarily the same as the

which are not necessatify the same as the target/epitope identified as the interactive species during the clinical stage of the auto-immune disease.

For example, a virus infection may trigger an immune reaction that yields a class of antibodies that cross-react with a normal cellular target. Such antibodies may bind more avidly to the original, stimulatory, viral antigen than to the epitope on the cellular target. As another example, the epitope on the cellular target may not take full advantage of the binding site on the antibody.

The polypeptide ligand is used diagnostically to measure the quantity of circulating auto-antibody, using, e.g., an ELISA assay. The technology is available to one skilled in the art, without undue experimentation. As another example, the fixed portion of the polypeptide ligand is used as the reporter substance when the polypeptide ligand interacts with the circulating auto-antibody. With a fixed carboxyterminus of beta-galactosidase or alkaline phosphatase, serum protein samples attached to plastic plates are assayed directly for the anti-H2B antibody by "staining" with the polypeptide ligand covalently fused (by recombinant DNA techniques) to either reporter enzyme.

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Example 3. Indirect Immunoprecipitation: Polypeptide ligands directed toward domains of any protein.

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Immunization of animals with antigens, whether crudely prepared or purified, often results in immune responses directed at a subset of the available epitomes in that antigen. The polyclonal sera may react largely with a single protein domain in that antigen. Similarly, when researchers attempt to raise antibodies against fusion proteins, often the well-known fusion partner is immuno-dominant over the new protein portion of the fusion.

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Antibodies aimed at a protein target (but that

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do not recognize the portion of the target that one wishes to use as the target in SPERT) allow INDIRECT Immunoprecipitation of ribosome complexes. That is, immunoprecipitation is a useful partitioning step when antibodies are aimed at domains in the target that are different from those domains pre-selected for SPERT-based ligand evolution. This protocol is sometimes called "panhandling", and can yield high-affinity polypeptide ligands for target domains that are weakly

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immunogenic.

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prepared as in Example 1 except that the randomized mRNA regions are now set to yield about 50 amino acids in the solvent-exposed nascent polypeptide. Biased randomization is done so that chain termination codons are not likely over the 150 randomized nucleotides; in addition, cell-free translation is performed in the presence of so-called suppressor tRNAs so that translation continues to the desired portion of the mRNAs.

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The population of ribosome complexes is pretreated with the antisera aimed at the target protein, but in the absence of that target protein. The pretreatment is designed to eliminate any nascent polypeptides that react directly with the antibodies, as in Example 1. The target protein is then added to the ribosome complexes, along with antibodies aimed at the target protein. Partitioning occurs as the ribosome complexes that interact with the target at the same time (see Figure 4).

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The single-stranded DNA binding protein of bacteriophage T4 (gp32) has an acidic carboxyterminal region which is immunodominant (K. Krassa, Ph.D., Thesis, 1987). In one immunization experiment, polyclonal sera react exclusively with the carboxyterminal domain of the protein; 12 monoclonal cell lines derived from hybridoma fusions with spleen

cells from such immunized animals produced antibodies that react with the same target domain. Purified polyclonal sera which react with the carboxy-terminal domain of gp32 are used for indirect immunoprecipitation in this example.

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A population of ribosome complexes is produced (above). These ribosome complexes are pre-treated with the polyclonal sera aimed at gp32; this is readily accomplished by passing the ribosome complexes through Staph A columns pre-bound with the polyclonal sera against gp32. Subsequently, those ribosome complexes unable to react directly with antibodies raised against gp32 are reacted with gp32, followed by treatment with the sera aimed at the carboxy-terminus of gp32. Goat anti-mouse antibodies are used to immunoprecipitate gp32 and whatever ribosomal complexes interact with the core domain of gp32. Cycles of SPERT are continued until a desired level of binding is attained. Sequences are then cloned and individuals identified and tested for affinity to gp32.

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Example 4. Isolation of a polypeptide ligand for a serine protease.

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Serine proteases are protein enzymes that catalyze hydrolysis of peptide bonds within proteins, often with high selectivity for specific protein targets (and, of course, for specific peptide bonds within the target protein). The serine proteases are members of a gene family in mammals. Examples of serine proteases are tissue plasminogen activator, trypsin, elastase, chymotrypsin, thrombin, and plasmin. Many disease states can be treated with polypeptide ligands that bind to serine proteases, for example, disorders of blood clotting. Elastase inhibitors are likely to be useful in minimizing the clinical progression of emphysema. Proteases other than serine

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proteases are also important in mammalian biology, and these too are targets for polypeptide ligands with appropriate affinities obtained according to the invention herein taught.

available for the binding of an inhibitory ligand. The Porcine elastase attached to agarose is available from identified and purified using the starting randomized buffer used for binding during the SPBRT cycles must material of Example 3. Serine proteases are easily check that the active site of the bound elastase is commercial sources. Thus, in this example affinity chromatography is the partitioning method. Natural elastase inhibitors are available, and are used to A ligand that binds to porcine elastase is dithiothreitol, which can reduce protein disulfide materials with retention of enzymatic activity. attached by standard methods to column support not denature or otherwise inactivate elastase; bonds, is left out of the binding buffer. ഗ 2 15

virtually all the nascent polypeptides are able to bind the column, the ribosome complexes are poured through a inhibitory ligand. The ribosome complexes that are not This procedure focuses the evolving polypeptide ligands of the mixture of nascent polypeptides becomes high, a rounds of SPERT are aimed at obtaining any polypeptide After several rounds of SPERT, as the affinity bound in this reversed elution procedure are used to inhibitory ligand for the elastase active site. In ligand that binds to any domain of elastase; after reversal of the elution parameters is used. Early prepare mRNAs for further SPERT cycles, once again depending on high affinity for the bound elastase. includes high concentrations of that same natural column that has been pre-saturated with a natural addition, the elution buffer for this procedure toward the elastase active site.

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When the mixture of polypeptide ligands has a

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demanding that the evolving polypeptide ligands have primarily toward the active site, further enrichment for high affinity inhibitors of elastase activity is accomplished by including low concentrations of the natural inhibitors in the partitioning steps, thus higher affinity than the effective affinity of the high affinity for the bound elastase, and is aimed natural inhibitor at the concentration used.

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ligands for other members of the serine protease family Nucleic acids encoding polypeptide ligands are in order to ascertain specificity within the family. efficiencies are measured with the same polypeptide cloned and seguenced, and binding affinities and inhibitory binding affinities for elastase are measured. Binding affinities and inhibitory

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receptor: A synthetic inhibitor of the Polypeptide ligands that antagonize a interleukin-1 receptor. Example 5.

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That domain serves as a binding site for cell extrinsic transduction through the membrane or internalization of either case polypeptide ligands of the invention may be molecules, including growth factors, peptide hormones, cause the normal activity of the natural ligand or to membrane such that a domain resides outside the cell. hormones), or even ions. Receptors handle the bound used to affect function of the receptor, that is to ligand in several different ways, including signal Receptors are a class of proteins that are the bound ligand for its subsequent function. In partially integrated into the cell's cytoplasmic non-peptide organic molecules (which may include block that activity.

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ligand through SPERT that is aimed at the interleukin-1 Receptor antagonism for a useful therapeutic purpose is accomplished by generating a polypeptide

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concentrations used in the work cited above. IL-lra is antagonist (called IL-lra for IL-1 receptor antagonist) (IL-1) receptor. A natural antagonist of the receptor competitive inhibitor of interleukin-1 binding to the (1990); Eisenberg et al., Nature, <u>343</u>:341-346 (1990), synthesized as a protein with 177 amino acids; after post-translational cleavage the active inhibitor has However, the activity of recombinant iL-lra, without glycosylation, is comparable to the activity of the 152 amino acids and, additionally, is glycosylated. receptor. The natural IL-1ra is a pure antagonist, has been found (Hannum et al., Nature, 343:336-340 and that antagonist has the presumptive utility of completely without agonist activity at the highest preventing or easing inflammatory problems such as those found in rheumatoid arthritis. The natural is partially homologous to IL-1 itself, and is a natural inhibitor.

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antagonist for the interleukin-1 receptor. Two methods principle recognize the features in common between IL-1 to the antigen combining site; such polypeptide ligands in Example 1, to develop polypeptide ligands that bind and IL-1ra. Those monoclonal antibodies are used, as Since one goal in this case is to provide antagonists are candidates for a novel class of IL-1 antagonists. SPERT is used to develop a polypeptide ligand raised against interleukin-1 that are able to crosspolypeptide is ca. 50 amino acids, as in Example 3. are used. In the first monoclonal antibodies are react with IL-1ra. Such monoclonal antibodies in smaller than the natural IL-1ra, the randomized

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the domain is attached to an insoluble matrix. complexes, are partitioned on the matrix. The matrix Candidate polypeptide ligands, residing in ribosome domain of the IL-1 receptor is itself used as the target for polypeptide ligand development through In a second methodology the extracellular 35

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polypeptides with the natural ligand known to bind to SPERT are continued until high affinity polypeptide the desired active site on the receptor. Cycles of is eluted with high concentrations of IL-1, thus displacing the ribosome complexes and nascent ligands are identified.

during SPERT that denatures the ribosome complexes even if the polypeptide ligand remains strongly bound to the Very high affinity, even covalent, antagonists protein denaturing conditions is used to prepare cDNA of the receptor are isolated by an elution protocol transcription provides mRNA for the next round of receptor. The mRNA eluted from the column under which is amplified through PCR, after which SPERT.

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and IL-1ra. Novel, SPERT-generated polypeptide ligands recognize structural or sequence homology between IL-1 IL-1 receptor antagonism. Those ligands identified by receptor-based affinity chromatography are tested with and characterized. SPERT-generated antagonists having novel antagonists recognized by those antibodies that having IL-1 receptor antagonist activity are isolated sequenced, and the polypeptide ligands are tested for less than 50% amino acid homology with natural IL-lra the antibodies of the first method to screen for the antagonists having less than 30% amino acid homology All genes encoding polypeptide ligands are are identified. In addition, SPERT-generated are identified.

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Example 6. Protein improvement by SPERT: Mutagenesis and selection of better natural insecticides.

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thuringiensis strains, have varying effectiveness for proteins. These proteins, derived from different B. Bacillus thuriengiensis is a gram-positive, spore-forming bacteria which produces insecticidal

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these insecticide proteins is to bind a receptor on the protein required to produce 50% mortality) can vary by one specific protein will kill the insect larvae of a killing insect larvae of different species. Although membranes serve as a functional partitioning tool in gut membranes of the susceptible insect larva. Such different insect targets (measured as the level of as much as 2000-fold. The mechanism of action for variety of species, the effectiveness toward the

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membranes from each of these insect larvae will be used suitable for SPERT by PCR; the appropriate DNA encodes which is fully active (Fischhoff et al., Biotechnology tomato hornworm and cabbage looper very effectively at insecticidal protein from t. subspecies kurstaki HD-1, required to kill tobacco budworm, corn earworm, black 5:807-813 (1987). This protein kills the larva of low concentration. Substantially more protein is cutworm, European cornborer, and beet armyworm. We create double-stranded DNA templates the N-terminal 646 amino acid portion of the as partitioning agents in SPERT.

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membranes is achieved. The DNA products are cloned and sequenced and individually assayed for effectiveness in binding membranes and larval killing. Effective toxins amino acid replacements. In particular, randomized DNA hypervariable region of the Bt. toxin. Rounds of SPERT ä are continued until a desired level of binding to gut sequence replaced by a sequence that is less than 50% The starting material in these experiments is methods are used to create protein variants. In one fixed codons within the insecticide, using about 50 are selected by SPERT, having a naturally-occurring throughout the 646 amino acids of the insecticide. RNA derived from the cloned gene, as above. Two method mutagenic PCR provides random mutations is used to replace the codons encoding the

toxic, SPERT-generated variants are identified wherein the original, naturally-occurring sequence is replaced by a sequence having less than 30% sequence homology In addition, homologous with the replaced sequence. with the replaced sequence.

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Inhibition of viral entry into target Anti-viral polypeptide ligands: Example 7.

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receptor. Rabies virus enters nerve cells through the acetylcholine receptor. Reovirus enters cells through is, host organisms can not easily alter such important epidermal growth factor receptor. Apparently viruses without suffering some impairment of crucial cell and Recently Kaner et al. (Science, <u>248</u>:1410-1413 virus, is said to enter cells through a cell adhesion the beta-adrenergic receptor. Vaccinia virus enters other viruses are said to utilize other receptors to through the CD4 glycoprotein receptor. Epstein-Barr (1990)) described the basic fibroblast growth factor Herpes Simplex Virus Type 1 (HSV) enters a cell. In Receptors are often used for viral attach on molecule ICAM-1. HIV, the AIDS virus, enters cells that same paper, by citation of other work several gain cellular entry. Rhinovirus, the common cold virus enters T lymphocytes via the C3d complement (FGF) receptor as the likely portal through which survive in part by using absolutely crucial cell receptors to gain entry into susceptible hosts. receptors so as to become resistant to the virus cells through a functional interaction with the organism functions. cells.

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identified that diminish viral uptake through receptors demonstrate a successful strategy. The soluble domain Polypeptide ligands of the invention are while still allowing critical growth factors to function. The basic FGF receptor is used to

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(1989)) is used as the target. A candidate mixture of of the basic FGF receptor (Lee et al., Science, 245:57 matrix bound extracellular domain of the FGF receptor. partitioning of ribosome complexes is obtained with polypeptide ligands is used as in Example 3.

ith high concentrations of FGF itself. Those ribosome complexes that remain on the column are further eluted polypeptides bind FGF receptors in a way that inhibits displaced by FGF contain nascent polypeptides that are The cycles of SPERT are altered to include an elution step from the matrix with high concentrations of HSV; Candidate polypeptides are assayed for during this elution step the ribosome complexes that avidly bound polypeptide that is eluted with FGF but HSV binding but does not interfere with FGF binding. exit the column are discarded, while those ribosome Several cycles of SPERT are used to find the most their negative impact on HSV infection and their candidate ligands with the desired specificity. complexes that are not displaced by HSV but are not with HSV. വ 10 15 20

neither antagonists nor agonists of the FGF receptor at polypeptides meeting these criteria are made using the criteria having less than 50% amino acid homology with the criteria having less than 30% homology with FGF is FGF is isolated. In addition, a polypeptide meeting most useful polypeptide ligands in this example are inability to prevent FGF-mediated cell growth. The concentrations that diminish HSV infection. Novel process as described. A polypeptide meeting the 22

Polypeptide ligands that enter cells: The glucocorticoid receptor and trojan horse ligands. Example 8.

isolated.

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steroid hormone, after which the receptor protein is The glucocorticoid receptor protein binds

internalized from the membrane so that the receptor can make its way into the cell nucleus. The receptor has a DNA binding domain (DBD) that interacts in the nucleus with target DNA sequences. Polypeptide ligands of the invention, agonists of the glucocorticoid receptor, are internalized along with the receptor, and thus directed sequentially to the cytoplasm and then to the nucleus. Depending on the dissociation rate constant for specific polypeptide ligands, these ligands largely reside after uptake in either the cytoplasm or the nucleus.

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ligands that enter cells are localized in the cytoplasm Example 3, SPERT is directed toward the glucocorticoid resistance to proteolysis of the polypeptide ligand is prior example, SPERT protocols are manipulated so that glucocorticoid binding domain but that have much lower abundance of the glucocorticoid receptor. Polypeptide the polypeptide ligands evolve, screening of potential ligands to provide cell entry for molecules with other affinity than that observed for steroid hormones. As receptor, either with indirect immunoprecipitation or or nucleus by means available to those skilled in the art. Those polypeptide ligands that enter cells with challenge, and testing both cells with and without an affinity chromatography using bound receptor. As in polypeptides are found that compete directly for the ligands is performed on individual candidates; thus tested using whole cell entry prior to the protease proper localization are fused to other polypeptide Using the randomized starting material of useful activities.

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Example 9. Polypeptide ligands toward nucleic acids: Inhibitors of transcription.

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Cancer cells can result from the over-expression of a transcriptional activator protein that

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functions to enhance transcription and subsequent expression of sets of genes that push the cell toward inappropriate and uncontrolled growth. Thus, mutations that elevate the activity of a transcriptional enhancer

of a set of genes relevant for growth control. Such tumors are treatable with polypeptide ligands that reset the appropriate level of expression or activity of the transcriptional enhancer. While it is likely that polypeptide ligands may be aimed at the enhancer protein directly, thus inhibiting the activity and resetting a proper growth rate, in the present example a polypeptide ligand is aimed at the production rate of the transcriptional enhancer.

The polypeptide ligand of interest binds to the genome of the cancer cell at a location that competes for transcription of the gene that encodes the transcriptional activator protein, and hence expression of that protein. That is, in classical genetic language, the polypeptide ligand is a specific transcriptional repressor.

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The starting materials of Example 3 are used to specific sequence of double-stranded DNA is prepared by chemical means and covalently attached to an insoluble ribosome complexes in general are able to flow through for all double-stranded DNA (that is, without adequate recovered, and placed into the SPERT protocol of mRNAcolumn matrix. The column matrix is chosen such that containing nascent polypeptide ligands that interact order to eliminate polypeptide ligands with affinity the column containing bound DNA. Ribosome complexes generate a mixed pool of candidate polypeptides. A amplification, transcription, and a second cycle. ribosome complexes are mixed with random soluble sequence specificity for the intended use), the specificity or not) are retarded on the column, with double-stranded DNA (either with sequence

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specific DNA sequences attached to the column. In this manner polypeptide ligands that are indifferent to DNA during the partitioning step than is the abundance of partitioning step. The soluble DNA concentration is adjusted to give about tenfold more non-specific DNA sequence emerge from the column along with ribosome louble-stranded DNA sequences prior to the column complexes containing polypeptide ligands that are unable to bind DNA at all.

second SPERT is directed toward the contiguous DNA base itself. This improvement in specificity and avidity is sequences are used to establish which nucleotide pairs polypeptide ligand with higher specificity and avidity in the covalently attached DNA are required for avid binding of the polypeptide (using the SELEX protocol described in U.S. Patent Serial No. 07/536,428). A than is available from either polypeptide ligand by polypeptide ligand, and the genes for the first and sequence are characterized further. Randomized DNA Polypeptide ligands aimed at a specific DNA second polypeptide ligands are combined to yield a containing a flexible peptide linker) to provide a "steps" are made independently and the polypeptide an example of walking, although in this case the polypeptide ligand fusion (in either order, and pairs that are not bound by the first isolated ligands joined post-identification.

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this example must overlap a transcriptional initiation The sequence of double-stranded DNA chosen in signal. The ras oncogene transcriptional initiation region is chosen first.

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Example 10. Human c-myc protein epitope.

select an epitope or epitopes from a random mixture of This experiment shows that it is feasible to RNA-encoded peptides. An antibody was chosen which

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recognizės an epitope in human c-myc protein consisting

translational initiation, insertion sites for random or of the amino acid sequence Glu-Gln-Lys-Iso-Ser-Glu-Glupromoter, a 5' untranslated region (5'-UTR) containing Asp-Lys (described in Evan et al., Mol. Cell. Biol. 5, 3610-3616, 1985). An expression system may be set up non-random sequences which would encode nascent for conducting SPERT experiments utilizing a T7 signals for either eukaryotic or prokaryotic

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peptides accessible to selection on ribosomes, and a 3' plasmid using oligos 9 and 10 from Table 3. These two ribosome. Refer to Table 3. The T7 promoter sequence oligos 1 and 2 from Table 3 using plasmid pSPBP4 which peptide sequences which are buried in the translating was added to the eukaryotic 5' UTR through PCR with is described by Siegel and Walter, (<u>Cell</u> 52: 39-49, 1988). The 3'-FTR was obtained by PCR of the same fixed translated sequence (3'-FTR) which encodes

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ligated. The ligated fragment was purified and further PCRd prior to cloning into the HindIII and BamHI sites of pBSSK+ (purchased from Strategene Systems, Inc.) to the HindIII and Nhe I site of pPSX-EUK to create pPSXwill be cloned using oligos 3 and 4 from Table 3 into create the plasmid pPSX-EUK. The prokaryotic 5'-UTR fragments, 5'-UTR and 3'-FTR were cut with NheI and

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with a prokarytic one. The myc epitope encoding insert PROK replacing the eurkaryotic ribosome binding site insert (for eight amino acids) is obtained by PCRing is obtained by PCRing the template oligo 7 with the oligos 5 and 6, all from Table 3, and the variable

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Table 3. These inserts will be digested with NheI and EcoRI and ligated in the presence of likewise digested insert in pPSX-EUK). Thus there will be a positive pPSX-EUK and pPSX-PROK. (This was done for the myc the template oligo 8 with the oligos 5 and 6, from control myc

epitope-encoding expression system which can be

the extent of translation using oligo 12 which puts two stop codons allowing repeated translation of individual translation with no amino acid depletion, and to test variable nascent peptide-encoding system which can be polysome complexes; this may be accomplished by using polysomes by the anti-myc antibody. Further testing separately by prokaryotic translation systems, and likewise variably translated, and a system with no oligos 10 in PCR (with oligo 1) to create multiple inserts which can serve as an internal control for will identify what 3' ends will give the stablest translated by eukaryotic translation systems and comparing extents of enrichments by selection of histidine codons for translation with no added histidine, with oligo 11 for normal unstopped

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WE CLAIM:

- A method for making a polypeptide ligand of a target molecule comprising:
 - synthesizing a translatable mRNA mixture comprising a ribosome binding site, translation initiation codon and a randomized sequence coding region; a)

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complexes, each member thereof comprising randomized coding region and said nascent polypeptide being the translation product a ribosome, a nascent polypeptide and a translated mRNA, said mRNA having a synthesizing a mixture of ribosome of said mRNA; â

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thereby separating the ribosome complexes partitioning the ribosome complexes with unbound complexes, the ribosome complexcomplexes to a desired target molecule, into ribosome complex-target pairs and target pairs having mRNA enriched for respect to binding of the ribosome sequences encoding target-binding polypeptides; ๋อ

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ribosome binding site, an initiation codon and a coding region enriched for sequences ribosome complex-target pairs to yield a translatable mRNA mixture comprising a encoding target-binding polypeptides; amplifying the mRNA of partitioned ਰ

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successive repeat as many times as desired to yield a desired level of target binding repeating steps b) through d) using the by a polypeptide encoded by the mRNA mRNA enriched for sequences encoding target-binding polypeptides of each enriched for sequences encoding the polypeptide; and ô

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 The method for selecting a polypeptide ligand of a desired target molecule from a polypeptide mixture comprising:

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a) synthesizing a polypeptide mixture each
 member thereof having attached thereto
 amplifying means for separately amplifying
 the individual polypeptide to which it is
 attached;

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b) partitioning the polypeptide mixture with respect to binding the target molecule, thereby separating the mixture into polypeptide-target pairs and unbound polypeptides;

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 c) amplifying the polypeptides of polypeptide-target pairs using said amplifying means; and

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- d) repeating the partitioning and amplifying steps to select a polypeptide ligand of a desired target molecule.
- The method of claim 2 wherein the polypeptide mixture comprises polypeptides having a segment of randomized amino acid sequence.

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4. The method of claim 3 wherein the segment of randomized amino acid sequence is from 4 to 50 amino acids in length.

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The method of claim 3 wherein the amplifying means comprises an mRNA mixture, each member thereof encoding a polypeptide of the polypeptide mixture and being attached to the polypeptide it encodes as part of a ribosome

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complex.

The method of claim 3 wherein the step of amplifying the polypeptides comprises the additional step of amplifying the mRNA mixture.

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The method of claim 6 wherein the mRNA mixture is amplified by reverse transcription and a polymerase chain reaction.

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- A method for making a polypeptide ligand of a target molecule comprising:
- (a) synthesizing a mRNA mixture comprising translatable and nontranslatable regions, wherein said translatable region comprises randomized and fixed sequence coding regions;

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(b) synthesizing a mixture of mRNA·polypeptide copolymers, each member comprising an mRNA and a polypeptide encoded by its associated mRNA, wherein a portion of said nontranslatable region of said mRNA and a portion of said polypeptide encoded by said fixed sequence coding region form a binding interaction;

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(c) partitioning the mRNA-polypeptide copolymers with respect to affinity of the copolymers to a desired target molecule;

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(d) amplifying the mRNA of partitioned copolymers to yield a translatable mRNA mixture; and

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- (e) synthesizing a polypeptide or polypeptides encoded by the mRNA mixture of step (d).
- 9. The method of claim 8 further comprising the steps of repeating steps (a) through (d) using the mRNA mixture of step (d) in successive

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		cycles repeating as many times as desired to yield copolymers with the desired affinity to the target.		21.	The method of claim 8 wherein the target molecule is a carbohydrate.
ĸ	10.	The method of claim 8 wherein the target molecule is a protein.	ហ	22.	The method of claim 8 wherein the target molecule is a polysaccharide.
		The method of claim 10 wherein the protein is an enzyme.		23.	The method of claim 8 wherein the target molecule is a lipid.
10	12.	The method of claim 10 wherein the protein is an antibody.	10	24.	The method of claim 8 wherein the target molecule is a glycolipid.
15	13.	The method of claim 10 wherein the protein is a receptor.	15	25.	The method of claim 8 wherein the target molecule is a toxin.
	14.	The method of claim 10 wherein the protein is a nucleic acid binding protein.		26.	The method of claim 8 wherein the target molecule is a drug.
20	15.	The method of claim 10 wherein the protein is a toxin.	20	27.	The method of claim 8 wherein the target molecule is a controlled substance.
	16.	The method of claim 10 wherein the protein is a glycoprotein.		28.	The method of claim 8 wherein the target molecule is a metabolite.
22	17.	The method of claim 10 wherein the protein is an antigen.	25	29.	The method of claim 8 wherein the target molecule is a cofactor.
30	18.	The method of claim 8 wherein the polypeptide is an inhibitor of function of the target molecule.	30	30.	The method of claim 8 wherein the target molecule is a nucleic acid.
	19.				molecule is a hormone.
35	20.	motecule is a cell membrane component. The method of claim 8 wherein the tarmet	32	32.	The method of claim 8 wherein the target molecule is a receptor ligand.
		molecule is a virus component.		33.	The method of claim 8 wherein the target

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bind to the bridging molecule anchored to the target molecules.

- polypeptide encoded by the second mRNA enriched The method of claim 8 comprising the additional steps a) - e) and a second randomized sequence using the second translatable mRNA mixture to yield a desired level of target binding by a mRNA mixture comprising the mRNA selected by steps of synthesizing a second translatable coding region, and repeating steps b) - e) for sequences encoding the polypeptide. 42.
- A mixture of mRNA polypeptide copolymers comprising: 43.

least a portion of the fixed sequence region of comprising random and fixed sequence regions, nontranslatable portion of said mRNA and at wherein said mRNA and polypeptide are bound a polypeptide encoded by said mRNA an mRNA comprising nontranslatable together by at least a portion of the portions and translatable portions; said polypeptide.

A polypeptide that is a ligand of a target molecule prepared according to the method described in claim 8.

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- A method for making a polypeptide ligand of target molecule comprising: 45.
- 1014 sequences comprising translatable and synthesizing a mRNA mixture of at least nontranslatable regions;
- copolymers, each member comprising an mRNA synthesizing a mixture of mRNA.polypeptide and a polypeptide encoded by its <u>a</u>

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associated mRNA, and not containing a ribosome;

- copolymers with respect to affinity of the copolymers to a desired target molecule; amplifying the mRNA of partitioned partitioning the mRNA.polypeptide છ g
 - synthesizing a polypeptide or polypeptides copolymers to yield a translatable mRNA mixture; and (e)
 - mRNA.polypeptide copolymers are synthesized by encoded by the mRNA mixture of step (d). nontranslatable portion of said mRNA and a the post-translational or co-translational interaction between a portion of the The method of claim 45 wherein said portion of said polypeptide.
- crosslinking the polypeptide-tRNA-mRNA complex mRNA polypeptide copolymers are synthesized by The method of claim 45 wherein said after translation of the mRNA. 47.
- mRNA to the initial amino acid sequences of the mRNA.polypeptide copolymers are synthesized by linking the 5' nucleic acid sequence of the The method of claim 45 wherein said polypeptide prior to translation.

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5. -TCCTGTCCATCGCAA-3'

\$ -455ADT65TA5D5TT65TADD- (-(50A) 021- }-55TA5D555- \$

5 - ACCATGGCAGCACTG-3'

'E-TOOTACOOTOOTOACOTO- [-NEA-]-OTTAAOOOO-'E

E-AADDDTDDDTDTDDTDDTADDDDDD-

'E-ADDADTƏDDADDƏTTƏDTADDƏTDƏTDADDTDƏTDTTATAADDƏD-'E

-DIATDIAAATDIDIATICOLOCITATATATATATTTTTAADDDD-'S

*E-AATAGAGAGATTTAGATAGAGGGGGAGATATGAGTGAGGATAATTGGAAGGG-*2

TABLE 1

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TABLE 2

1.1 5' fixed sequence

Hind!!! site

Ribosome binding site EcoRI

5'-CCGAACCTTAATACGACTCACTATAGGGCGACATACATTTACACATAAggaggaauuauaauatgaaagaalteecg-3'
3'-99ciicgaallalgecgaaglgalaleeegelGTATGTAAATGTGTGTATTCCTCCTTATATATATATTTCTTAAGGGC-5'

- T7 promoter

2.1 Stratagene polylinker cloning site (pBSSK+)

Pati

S'-TCGATAAGCTTCATATCGAATTCCTGCAGCCCGGGGGATCCACTAG-3'

ECORI

Hindii

3.) 3' primer annealing aite and insertion sequence cloning sites

NCOI Pati Ecori

BAMI

4.) Randomizing oligonuclectides to be cloned at the EcoRi, Pati, and Ncol sites.

PotI

A./ 5'-cccGAATTC-[-45N-]-CTGCAGTGCTGCCATGGT-3'

NC01 NCOI 9.) s'-666CCATGG-[-120 (ACG)-1-CCATGGTTGCGATGGTCAGGA-3' 3' -AACGCTACCTGTCCT-5'

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TABLE 3

1. PE5 (5' primer for 5' untranslated region (UTR) and full-length PCR)

5'-GGGAAGCTTAATACGACTCACTATAGGGAGCTTGTTCTTTTTGCAGAAGCTCAG-3'

2. 3'UTR (3' Primer for PCRing the 5' untranslated region prior to ligation)

5'-crescetascearsstarterscearasticas-3'

3. PROTOP (5' primer for fixed proke UIR-RBS PCR and cloning)

4. PROBOT (Complement to Protop for cloning proke UTR-RBS)

5' - CIAGCCATITIAITITCCTCCTIAICTIACCTIACCCTAIAGTGAGTCGTAITAAGCTTGGG -3'

5'-GGGCCATGGCTAGCGCCGAGGA-3'

PH3 (3' primer for fixed epitope (EPI) and variable region (VAR) PCR, sequencing and (maybe) cloning)

5'-GGCGGATCCAGGCGGACCCTTTCTGCGACGAA-3'

7. MycCODE (oligo for EPI construction)

5'-cGGCCAIGGCIAGCCCGAGGAGCAGAAGCIGATCICCGAGGAGCTGCTGGAAATICGICGCAGAAAGGGTCCCG-3'

8. VarCODE (oligo for VAR construction)

5'-GGGCCATGGCTAGCGCCGAGGAGHNNNNNNNNNNNNNNNNNNNNNNNNNCTGCTGGAATTCGTCGCAGAAAGGGTCCCG-3'

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3'fixed 113'primer 1 sequence 11 anneal-1 ing site

| randomized sequence

rototini codon

mRNA | ribosome binding

IN VITRO TRANSLATION

ribosome

translated_mRNA

SELECTION

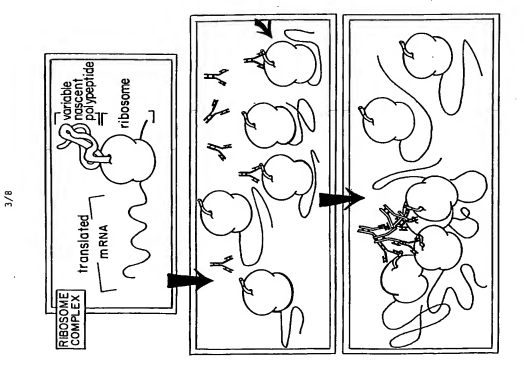
Ribosome complex

variabe Discent

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BOUND

F1G. 1



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F16.3

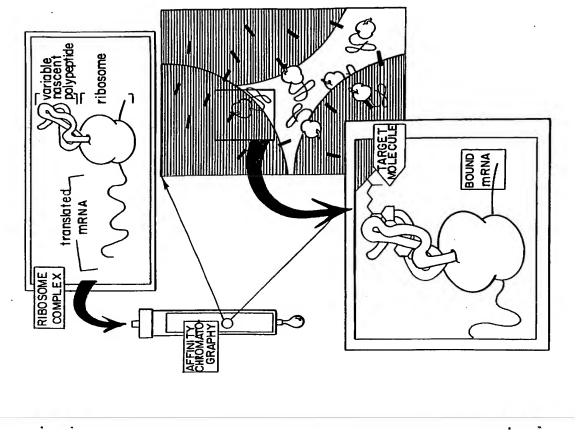
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ribosome

translated

RIBOSOME COMPLEX mRNA



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F16.5

ATTACHED GUIDE EPI TOPE

INHIBITOR

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GUIDE EPITOPE ANCHORED AT THE TARGET SITE.

> GUIDE INTERACTION

PROTEIN OF INTERESIN

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SECONDARY INTER ACTION

INTERNATIONAL SEARCH REPORT

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CLASSIFICATION OF SUBJECT MATTER

5) :CI2P 21/00; CI2N 15/09; C07H 21/02; C07K 7/00, 13/00, 15/00

CL. :Perio See Extra Shea.

secording to International Patent Classification (IPC) or to both national classification and IPC

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Minimum documentation searched (classification system followed by classification symbols)

U.S. : Please See Extra Sheet.

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronio data base consulted during the international search (name of data base and, where practicable, search terms used)

Piease See Extra Sheet.

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Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
>	European Journal of Biochemistry, Volume 112, issued 1980, A.J.M. Wagenmakers et al, "Cross-linking of mRNA to Proteins by Irradiation of Intact Cells with Ultraviolet Light", pages 323-330, entire document.	43, 45-48
×	Nature, Volume 343, issued 25 Innuary 1990, S. Eisenberg et al. "Primary Structure and functional Expression from complementary DNA of a human interleukin-1 receptor Anagonist", pages 341-346, entire document.	4
<	GB, A, 2,183,661 (Ballivet et al) 10 June 1987, entire document.	148
×	Science, Volume 249, issued 27 July 1990, J. Scott et al. Searching for Peptide Ligands with an Epitope Library. pages 386-390, entire document	44 1-43, 45-48
××	WO, A, 91/05038 (Kawasaki) 18 April 1991, entire document.	1-7, 44 8-43, 45-48

See patent family annex.	T bater document published after the internation
Further documents are listed in the continuation of Box C.	Special categories of cited documents:
	۰

 Annual published after the international filing date or priority drawers not not in conflict with the application has the file of understand the principle or theory underlying the invention document of particular relevance; the claimed invention cannot be associated to inventorle as invention sup. when the document a combined with one or more other such documents, such combination being obvirous to a person skilled in the art document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone document member of the same patent family × . document defining the general state of the 2st which is not considered to be part of particular relevance document which may throw deaths an priority chimits) or which is eited to establish the publication date of usother chains or other special reason (as specified) document referring to an and disclosure, use, exhibition or other means earlier document published on or after the international filing date

Date of mailing of the international search report document published prior to the international filing date but later than $~~\cdot_{K^*}$ the priority data claimed Date of the actual completion of the international search

GABRIELE E. BUGAISKY Telephone No. (703) 308-0196 Authorized officer Name and mailing address of the ISA/ Commissioner of Patents and Trademarks Box PCT Weshington, D.C. 20231 Facsimile No. NOT APPLICABLE 16 September 1992

orm PCT/ISA/210 (second sheet)(July 1992)*

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INTERNATIONAL SEARCH REPORT

International application No. PCT/US92/00801

A. CLASSIFICATION OF SUBJECT MATTER:

US CL:

435/68.1, 69.1, 69.2, 69.3, 69.4, 69.5, 69.6, 172.1; 530/300, 350; 536/27

B. FIELDS SEARCHED

435/68.1, 69.1, 69.2, 69.3, 69.4, 69.5, 69.6, 172.1, 172.3; 530/300, 350; 536/27; 935/3, 4, 10, 11, 12, 13, 14, 15, Minimum documentation scarched Classification System: U.S.

B. FIELDS SEARCHED

16, 17, 21, 44,

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, DIALOG files 155, 5, 73, 357, search terms: reverse translation, polymerase chain reaction, cell free translation, in vitta translation, ligand, bind, directed evolution, transcription translation system, transcription, translation, system, RNA, mRNA, polypeptide(s), peptide(s), protein(s), covalient, polymer(s), copolymer(s), copolymer(s)

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